WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT) (51) International Patent Classification 5: WO 93/12151 (11) International Publication Number: A1 C08F 210/02, 4/60, 210/16 (43) International Publication Date: 24 June 1993 (24.06.93) PCT/US92/10590 (74) Agents: BELL, Catherine, L. et al.; Exxon Chemical Com-(21) International Application Number: pany, P.O. Box 2149, Baytown, TX 77522-2149 (US). (22) International Filing Date: 11 December 1992 (11.12.92) (81) Designated States: CA, JP, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, (30) Priority data: 806,894 13 December 1991 (13.12.91) US (71) Applicant: EXXON CHEMICAL PATENTS INC. [US/ **Published** US]; 5200 Bayway Drive, Baytown, TX 77520-5200 (US). With international search report. (72) Inventors: BRANT, Patrick; 103 Harborcrest, Seabrook, TX 77586 (US). CANICH, Jo, Ann, Marie; 900 Henderson Avenue, Apt. 808, Houston, TX 77058 (US).

(54) Title: ETHYLENE/LONGER ALPHA-OLEFIN COPOLYMERS

(57) Abstract

High molecular weight linear copolymers of ethylene and 1-50 mole percent linear α -olefins having from 10 to 100 carbon atoms are disclosed. The polymers have M_w of 30,000-1,000,000, MWD of 2-4, a density of 0.85-0.95 g/cm³, and a high composition distribution breadth index. Also disclosed are a method for making the polymers with a cyclopentadienyl metallocene catalyst system, and adhesives, films, molded articles and other products made from the copolymers.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

ATU BB BE BF BG BJ BR CA CF CG CM CZ DE ES FI	Austria Australia Barbados Belgium Burkina Faso Bulgaria Benin Brazil Canada Central African Republic Congo Switzerland Cöte d'Ivoire Cameroon Czechoslovakia Czech Republic Germany Denmark Spain Finland	FR GA GB GN GR HU IE IT JP KP KR LI LK LU MC MC MI MN	France Gabon United Kingdom Guinea Greece Hungary treland Italy Japan Democratic People's Republic of Korea Republic of Korea Kazakhstan Licehtenstein Sri Lanka Luxembourg Monaco Madagasear Mali Mongolia	MR MW NL NO NZ PL PT RO RU SD SE SK SN TD TG UA US VN	Mauritania Malawi Netherlands Norway New Zealand Poland Portugal Romania Russian Federation Sudan Sweden Slovak Republic Senegal Soviet Union Chad Tugo Ukraine United States of America Viet Nam
---	--	---	---	---	---

WO 93/12151 PCT/US92/10590

1

ETHYLENE/ LONGER ALPHA-OLEFIN COPOLYMERS Field of the Invention

This invention relates to copolymers of ethylene and longer α -olefins. More particularly, this invention relates to high molecular weight ethylene copolymers of α -olefins having from 10 to 100 carbon atoms. This invention also relates to a process for copolymerizing ethylene with longer α -olefins utilizing certain transition metal compounds from Group IV B of the Periodic Table of Elements that produces high molecular weight copolymers.

10 Background of the Invention

30

35

Prior art copolymers of ethylene and longer α -olefins, i.e. olefins having about 10 or more carbon atoms, have suffered a number of disadvantages. Primarily, it has not possible to prepare such copolymers, 15 sufficiently high molecular weight for most applications, using a traditional Ziegler-Natta catalyst. Generally, the longer α -olefin comonomer content increases, the molecular weight decreases significantly. Also, ethylene copolymers have had a very broad molecular weight 20 distribution, as well as a broad α -olefin composition distribution. This arises from the prior art catalyst systems having a high ratio of ethylene:comonomer reactivity, as well as a low ratio of polymerization propagation to termination. As a result, the low molecular weight species have substantially higher α -olefin comonomer content and the high molecular weight species have a very low comonomer content.

It has been proposed to use certain metallocenes such as bis(cyclopentadienyl) titanium or zirconium dialkyls in combination with aluminum alkyl/water cocatalyst as a homogeneous catalyst system for the polymerization of olefins. For example: German Patent Application 2,608,863 teaches the use of a catalyst system for the polymerization of ethylene consisting of bis(cyclopentadienyl) titanium dialkyl, aluminum trialkyl and water: German Patent Application 2,608,933 teaches an ethylene polymerization catalyst system consisting of zirconium metallocenes of the

35

general formula (cyclopentadienyl),ZrY4-n, wherein n stands for an integer in the range of 1 to 4, Y for R, CH2AlR2, $CH_2CH_2AlR_2$ and $CH_2CH(AlR_2)_2$, wherein R stands for alkyl or metallo alkyl, an aluminum trialkyl cocatalyst and water; European Patent Application No. 0035242 teaches a process for preparing ethylene and atactic propylene polymers in the presence of a halogen-free Ziegler catalyst system of (1) formula of cyclopentadienyl compound $(cyclopentadienyl)_{n}MY_{4-n}$ in which n is an integer from 1 to 4, M is a transition metal, especially zirconium, and Y is 10 either hydrogen, a C_1 - C_5 alkyl or metallo alkyl group or a formula CH2AlR2, radical having the following general $CH_2CH_2AlR_2$, and $CH_2CH(AlR_2)_2$ in which R represents a C_1-C_5 alkyl or metallo alkyl group, and (2) an alumoxane; and U. Patent 4,564,647 teaches a low pressure process for 15 polymerizing ethylene, either alone or in combination with small amounts of other lpha-olefins, in the presence of a catalyst which may comprise a cyclopentadienyl compound, wherein (Cp) MR²R³R⁴ formula the represented by represents a cyclopentadienyl group, M represents titanium, 20 vanadium, zirconium or hafnium, and \mathbb{R}^2 , \mathbb{R}^3 and \mathbb{R}^4 are each an carbon atoms, to 6 group having from 1 cyclopentadienyl group, a halogen atom or a hydrogen atom, an alumoxane, which can be prepared by reacting trialkyl aluminum or dialkyl aluminum monohalide with water and a 25 Each of the above patents also teach that the polymerization process employing the homogeneous catalyst system is hydrogen sensitive thereby providing a means to control polymer molecular weight.

As is well known in the prior art, catalyst systems comprising a cyclopentadienyl compound, hereinafter frequently referred to as a metallocene or metallocene catalyst component, and an alumoxane offer several distinct advantages when compared to the more conventional Ziegler-type catalyst systems. For example, the cyclopentadienyl-transition metal/alumoxane catalyst systems, particularly those wherein the cyclopentadienyl compound contains at least one halogen atom, have demonstrated extremely high activity in the polymerization of α -olefins, particularly

WO 93/12151 PCT/US92/10590

4

10

15

3

ethylene. Moreover, these catalyst systems produce relatively high yields of polymer product having a relatively narrow molecular weight distribution. However, these catalyst systems, when used to prepare copolymers of ethylene with longer α -olefins in anything more than a very minor proportion, still suffer from the drawbacks of low incorporation rates, and low molecular weights.

For many applications it is of primary importance for a polyolefin to have a high weight average molecular weight relatively while having a narrow molecular distribution. A high weight average molecular weight, when accompanied by a narrow molecular weight distribution, polyolefin or an ethylene-lower-α-olefin copolymer with high strength properties. Traditional Ziegler-Natta catalyst systems --a transition metal compound cocatalyzed by an aluminum alkyl -- are capable of producing polyolefins having a high molecular weight but a broad molecular weight distribution.

More recently a catalyst system has been developed 20 wherein the transition metal compound has two or more cyclopentadienyl ring ligands, such transition metal compound also being referred to as a metallocene -- which catalyzes the production of olefin monomers to polyolefins. Accordingly, metallocene compounds of the Group IV B metals, particularly, titanocene and zirconocene, have been utilized 25 as the transition metal component in such "metallocene" containing catalyst system for the production of polyolefins and ethylene- α -olefin copolymers. When such metallocenes are cocatalyzed with an aluminum alkyl -- as is the case 30 with a traditional type Ziegler-Natta catalyst system -- the catalytic activity of such metallocene catalyst system is generally too low to be of any commercial interest. since become known that such metallocenes may be cocatalyzed with an alumoxane -- rather than an aluminum alkyl -- to provide a metallocene catalyst system of high activity which 35 catalyzes the production of polyolefins. The zirconium metallocene species, as cocatalyzed or activated with alumoxane are commonly more active than their hafnium or

titanium analogues for the polymerization of ethylene alone or together with a lower α -olefin comonomer.

A wide variety of Group IV B transition metal compounds of the metallocene type have been named as possible candidates for an alumoxane cocatalyzed catalyst system. Hence, although bis(cyclopentadienyl) Group IV B transition metal compounds have been the most preferred and heavily use in metallocenes for type investigated metallocene/alumoxane catalyst for polyolefin production, that appeared have 10 suggestions tris(cyclopentadienyl) transition metal compounds may also be useful. See, for example, U. S. Patents Nos. 4,522,982; Such mono(cyclopentadienyl) .4,530,914 and 4,701,431. transition metal compounds as have heretofore been suggested as candidates for a metallocene/alumoxane catalyst are mono(cyclopentadienyl) transition metal trihalides and trialkyls.

15

25

30

35

More recently International Publication No. WO 87/03887 described the use of a composition comprising a transition metal coordinated to at least one cyclopentadienyl and at least one heteroatom ligand as a metallocene type component for use in a metallocene/alumoxane catalyst system for α olefin polymerization. The composition is broadly defined as a transition metal, preferably of Group IV B of the Periodic Table which is coordinated with at least one cyclopentadienyl ligand and one to three heteroatom ligands, the balance of the coordination requirement being satisfied The hydrocarbyl ligands. cyclopentadienyl or described is system catalyst metallocene/alumoxane illustrated solely with reference to transition metal compounds which are bis(cyclopentadienyl) Group IV B transition metal compounds.

Therefore, a need still exists for catalyst systems that permit the production of higher molecular weight ethylene-longer- α -olefin copolymers and desirably with a molecular weight distribution narrow and The present invention addresses composition distribution. the need, then, for a polymerization process which permits the efficient and economically attractive production of high

WO 93/12151 PCT/US92/10590

5

molecular weight ethylene/longer α -olefin copolymers and copolymer products.

Summary of the Invention

10

30

In accordance with the present invention, longer α olefins are copolymerized with ethylene in the presence of a catalyst system comprising an activated cyclopentadienyltransition metal compound. Quite surprisingly, it has been found that the longer α -olefins have a polymerization rate on the same order as ethylene when these catalysts are employed, despite the large "tail" of the longer α -olefin. As result, the longer α -olefin is unexpectedly incorporated into the copolymer at a competitive rate with the ethylene, and the composition distribution substantially uniform and random.

15 The present invention resides, at least in part, in the discovery that longer α -olefins (e.g. C₁₀-C₁₀₀) polymerized with ethylene using certain monocyclopentadienyl metallocene catalysts to obtain a high molecular weight copolymer with a high proportion of longer α -olefin 20 incorporation, a narrow molecular weight distribution and a relatively random and uniform longer α -olefin comonomer distribution. Certain of these copolymers have very surprising properties, such as, for example, modulus, strain to break, rheological properties, storage and loss moduli, dissipative characteristics, and the like, as detailed more 25 completely below.

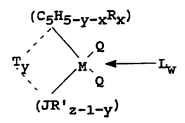
In one aspect, then, the present invention provides a substantially compositionally uniform copolymer of ethylene and from about 1 to about 50 mole percent, preferably from about 2 to about 30, and especially from about 4 to about 30 mole percent, of a longer α -olefin having at least 10 carbon atoms, preferably at least 12 carbon atoms. The copolymer has a density of from about 0.85 to about 0.95 g/cm³, and can be semicrystalline or amorphous. The copolymer preferably has a weight average molecular weight from about 30,000 to about 1,000,000 daltons or more, more preferably from about 80,000 to about 500,000 daltons, and a molecular weight distribution substantially between about 2 and about

20

4. The copolymer has a generally uniform comonomer composition distribution.

In another aspect, the present invention provides adhesives comprising a blend of the foregoing copolymer with a tackifier. The present invention also provides useful articles made from the foregoing copolymers, including films, sheets, coatings and molded articles.

In a further aspect, the present invention provides a method of preparing a copolymer by contacting ethylene and α -olefin having at least 10 carbon atoms with a catalyst at polymerization conditions wherein the ethylene:comonomer reactivity ratio is less than about 50. In a preferred embodiment, the foregoing copolymers are prepared by contacting ethylene and a longer α -olefin with a catalyst system comprising an activated Group IV B transition metal component at polymerization conditions, and recovering a high molecular weight, narrow molecular weight copolymer having a generally uniform, random α -olefin composition distribution. The "Group IV B transition metal component" of the catalyst system is represented by the general formula:



wherein: M is Zr, Hf or Ti and is in its highest formal oxidation state (+4, d^0 complex);

 $(C_5H_{5-y-x}R_x)$ is a cyclopentadienyl ring which is substituted with from zero to five substituent groups R, "x" is 0, 1, 2, 3, 4 or 5 denoting the degree of substitution, and each substituent group R is, independently, a radical selected from a group consisting of C_1-C_{20} hydrocarbyl radicals, substituted C_1-C_{20} hydrocarbyl radicals wherein one or more hydrogen atoms is replaced by a halogen radical, an amido radical, a phosphido radical, an alkoxy radical, an alkylborido radical, or any other radical containing a Lewis

15

20

25

30

35

acidic or basic functionality, C_1-C_{20} hydrocarbyl-substituted metalloid radicals wherein the metalloid is selected from the Group IV A of the Periodic Table of Elements, and halogen radicals, amido radicals, phosphido radicals, alkoxy radicals, alkylborido radicals or any other radical containing Lewis acidic or basic functionality or $(C_5H_{5-y-x}R_x)$ is a cyclopentadienyl ring in which two adjacent R-groups are joined forming a C_4-C_{20} ring to give a saturated or unsaturated polycyclic cyclopentadienyl ligand such as indenyl, tetrahydroindenyl, fluorenyl or octahydrofluorenyl;

 (JR'_{z-1-y}) is a heteroatom ligand in which J is an element with a coordination number of three from Group V A or an element with a coordination number of two from Group VI A of the Periodic Table of Elements, preferably nitrogen, phosphorus, oxygen or sulfur, and each R' is, independently a radical selected from the group consisting of C_1-C_{20} hydrocarbyl radicals, substituted C_1-C_{20} hydrocarbyl radicals wherein one or more hydrogen atoms is replaced by a halogen radical, an amido radical, a phosphido radical, an alkoxy radical, an alkylborido radical or any other radical containing a Lewis acidic or basic functionality and "z" is the coordination number of the element J;

Each Q may be independently any univalent anionic ligand such as halogen, hydride, or substituted or unsubstituted C_1 - C_{20} hydrocarbyl, alkoxide, aryloxide, amide, arylamide, phosphide or arylphosphide, provided that where any Q is a hydrocarbyl such Q is different from $(C_5H_{5-y-x}R_x)$ or both Q together may be an alkylidene or a cyclometallated hydrocarbyl or any other divalent anionic chelating ligand;

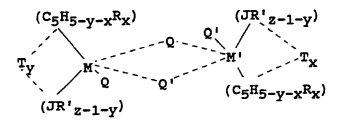
"y" is 0 or 1 when w is greater than 0; y is 1 when w is 0; when "y" is 1, T is a covalent bridging group containing a Group IV A or V A element such as, but not limited to, a dialkyl, alkylaryl or diaryl silicon or germanium radical, alkyl or aryl phosphine or amine radical, or a hydrocarbyl radical such as methylene, ethylene and the like;

L is a Lewis base such as diethylether, tetraethylammonium chloride, tetrahydrofuran, dimethylaniline, aniline, trimethylphosphine, n-butylamine,

15

20

and the like; and "w" is a number from 0 to 3; L can also be a second transition metal compound of the same type such that the two metal centers M and M' are bridged by Q and Q', wherein M' has the same meaning as M and Q' has the same meaning as Q. Such compounds are represented by the formula:



The metallocene catalyst component can be activated by an alumoxane component represented by the formulas: $(R^3-Al-0)_m$; $R^4(R^5-Al-0)_m-AlR^6_2$ or mixtures thereof, wherein R^3-R^6 are, independently, a univalent anionic ligand such as a C_1-C_5 alkyl group or halide and "m" is an integer ranging from 1 to abut 50 and preferably is from about 13 to about 25. Alternatively, the metallocene catalyst component can be activated with a cation capable of donating a proton and a bulky, non-coordinating anion capable of stabilizing the metal cation formed by reaction between the proton provided by the cation and a substituent of the metallocene reactive with the proton.

Brief Description of the Drawings

Fig. 1 is a semilog plot of loss tangent versus temperature for an adhesive of the present invention comprising a 60:40 weight blend of ethylene/hexadecene copolymer and ESC-5380 tackifier (Example 23) showing glass transition temperature.

Fig. 2 is a semilog plot of loss tangent versus temperature for an adhesive comprising a 60:40 blend of ethylene/hexadecene copolymer and ESC-1310LC tackifier showing glass transition temperature.

Fig. 3 is a log-log plot of viscosity ($\Delta-\Delta-\Delta$) and stress ($\diamond-\diamond-\diamond$) at 75°C (steady flow) versus frequency for

20

25

30

an ethylene/octadecene copolymer (Example 21) of the present invention.

Fig. 4 is a plot of the same material and variables of Fig. 3 at 140° C.

Fig. 5 is a log-log plot of viscosity $(\Delta-\Delta-\Delta)$ and stress $(\diamondsuit-\diamondsuit-\diamondsuit)$ at 25°C (steady flow) versus frequency for an ethylene/dodecene copolymer (Example 14).

Fig. 6 is the plot of the same material and variables of Fig. 5 at 95°C.

Fig. 7 is the plot of the same material and variables of Fig. 5 at 75° C.

Fig. 8 is a semilog plot of tan δ (10% strain, G" (100 rad/sec)/G'(1 rad/sec)) versus mole percent dodecene (Examples 8 and 10) and tetradecene (Examples 13 and 14) comonomer in the ethylene copolymers of the present invention compared to ethylene/butene copolymers.

Fig. 9 is a log-log plot of storage modulus (G') versus frequency for an ethylene/tetradecene copolymer $(\Delta-\Delta-\Delta)$ (Example 14) and an ethylene/octadecene copolymer $(\diamondsuit-\diamondsuit-\diamondsuit)$ (Example 21) compared to ethylene/propylene and ethylene/butene copolymers.

Fig. 10 is a plot showing the deviation in weight average molecular weight (M_) from standard polyethylene calibration curve (ratio of M, determined by viscometric GPC to M, determined by GPC differential refractive index (DRI) using polyethylene calibration M_(GPC/VIS)/M_(GPC/DRI PE)) versus mole percent comonomer in ethylene/α-olefin copolymers for comonomers propylene, butene-1, hexene-1, dodecene-1, tetradecene-1 and octadecene-1.

Fig. 11 is a plot of correction factors for converting observed molecular weight into actual molecular weight for the copolymers of the present invention.

Detailed Description of the Invention

35 The present invention relates to copolymers of ethylene with longer α -olefins. The longer α -olefins are preferably linear monomers of at least 10 carbon atoms up to about 100 carbon atoms or more. The novel characteristics of the

15

20

25

30

35

copolymers of the present invention derive from the side chains that are long pendant alkyl relatively introduced by the "tails" of the longer lpha-olefins comonomers as they are inserted into the generally linear polymer chain. When the side chains reach about 8 carbons in length (corresponding to decene-1 comonomer), and the side chains are sufficiently prevalent in the polymer, the side chains imparting capable of crystallization and characteristics to the polymer. Particularly at side chain lengths of 10 or more carbon atoms (corresponding to C_{12} α olefin comonomer), the crystallizability of the side chains Theoretically, any α is more definite and pronounced. olefin up to 100 carbon atoms or more is used to impart side chain crystallizability, but as a practical matter, α olefins of up to C_{30} of the desired purity are available commercially. Alpha-olefin monomers having more than about 30 carbon atoms generally have a broader distribution of molecular weights, and can also have some branching which influences crystallizability. Thus, the preferred α -olefins in this invention are linear α -olefins having from about 10 to about 100 carbon atoms, more preferably from about 12 to about 30 carbon atoms.

Specific representative examples of the longer 1-dodecene, 1-decene, 1-undecene, include olefins 1-hexadecene, 1-octadecene, 1-eicosene, tetradecene, docosene, 1-tetracosene, 1-hexacosene, 1-octacosene, 1-1-tetracontene, 1-dotriacontene, triacontene, pentacontene, 1-hexacontene, 1-heptacontene, 1-octacontene, 1-nonacontene, 1-hectene and the like. In general, the longer the α -olefin, the more pronounced are the properties as the size of the α -olefin imparted thereby, e.g. increases, the more unlike polyethylene the copolymer As the size of the comonomer increases, the becomes. softness, for example, generally increases while strain to a point where break decreases, up to crystallinity occurs, and then, quite surprisingly, softness decreases with additional comonomer length and strain to The copolymer can further contain break increases. additional monomers usually in relatively minor amounts,

WO 93/12151 PCT/US92/10590

10

15

20

25

30

35

11

which do not substantially adversely affect the novel Such termonomers include properties of the copolymers. vinyl and vinylidene compounds, for example, lower α -olefins having from 3 to 9 carbon atoms, such as propylene, 1-pentene, 3-methyl-pentene-1, butene. isobutene, methylpentene-1. 1-hexene, 1-heptene, 1-octene, 3,3,5trimethylpentene-1, 1-nonene, vinyl cyclohexene, and the like; dienes, such as 1,3-butadiene, 1,5-hexadiene and the like; vinyl aromatic monomers, such as styrene or alkylsubstituted styrene and the like; and combinations thereof.

Preferably, the ethylene is interpolymerized with from about 1 to about 50 mole percent longer α -olefin, more preferably from about 2 to about 30 mole percent longer α olefin, and especially from about 4 to about 30 mole percent longer α -olefin. In general, at an increased longer α olefin content, the properties imparted by the longer α olefin are more pronounced, e.g., density and strain to break decrease while softness increases with increasing longer α -olefin content. However, when the α -olefincomonomer content is increased to a point where the side chains become prevalent, e.g. there is side crystallization as a separate domain, the softness decreases and strain to break increases significantly.

The polymers of the present invention can vary from completely to semicrystalline. The amorphous ethylene/longer α -olefin copolymers generally have a density from about 0.85 to about 0.95 g/cm3. Amorphous material density generally has a below about 0.87 Semicrystalline polymer is generally in the density range of from about 0.854 to about 0.92 g/cm3. Crystallinity can be influenced by a number of factors, including molecular weight, the size of the longer α -olefin and content thereof, and the composition distribution. In general, copolymers containing about 12 mole percent or more of randomly distributed longer α-olefin are amorphous, copolymers containing less than about 12 mole percent of the longer α -olefin comonomer have more crystallinity as the comonomer content is reduced.

15

the present invention have of polymers surprisingly high molecular weight, preferably from about 30,000 to about 1,000,000 daltons or more, depending on the As used herein, molecular desired end-use application. weight refers to the weight average molecular weight (M_w) , unless otherwise indicated. The unique characteristics of the longer α -olefin copolymers are not generally observed at lower molecular weights where there is limited chain entanglement. Polymers having a molecular weight higher than this range, while theoretically possible, are difficult to prepare as a practical matter. Most commercially useful polymers, e.g. in film and adhesive applications, have M in the range of from about 80,000 to about 500,000 daltons.

The polymers of the present invention have a narrow molecular weight distribution (MWD). This surprising fact is reflected in a low polydispersity, i.e. a ratio of M_{ν} to number average molecular weight (M_{ν}) . The MWD (M_{ν}/M_{ν}) is generally in the range of from about 2 to about 4, even in the copolymers of very high molecular weight.

invention are present copolymers of the 20 substantially random and quite surprisingly have a fairly α -olefin distribution throughout uniform longer This uniform composition is reflected in a copolymer. relatively high composition distribution breadth index As used herein, CDBI is defined as the percentage 25 by weight of the copolymer molecules having a longer α olefin comonomer content within 50 percent of the median molar comonomer content, i.e. ± 50 percent of the median C_{10} -Homopolymers such as polyethylene, C₁₀₀ olefin content. which do not contain a comonomer, thus have a CDBI of 100%. The CDBI of a copolymer is readily calculated from data obtained by techniques known in the art, such as, example, temperature rising elution fractionation (TREF) as described in U. S. Ser. No. 151,350 or Wild et al., J. Poly. Sci. Poly. Phys. Ed., vol. 20, p. 441 (1982). ethylene/longer α -olefin copolymers herein generally have a CDBI on the order of about 70 percent or more, i.e. about 70 percent or more of the copolymer has a molar longer α -olefin comonomer content within ±50 percent of the median comonomer

WO 93/12151 PCT/US92/10590

s

15

20

25

30

35

13

content. In contrast, linear low density polyethylene prepared using conventional Ziegler-Natta catalyst has a CDBI on the order of 30 to 40 percent.

The present polymers comprise linear, comb-like molecules, as opposed to uncontrolled long chain branched polymers. This derives from the use of a single-site coordination catalyst as opposed to a free radical catalyst. The olefin polymerizes in a predominantly head-to-tail fashion so that the polymer molecule has a generally linear main chain formed by polymerization at the carbon-carbon double bond, and a plurality of side chains of controlled length corresponding to the aliphatic "tails" of the longer α-olefin.

The novel characteristics of the ethylene/longer α -olefin copolymers of the present invention, i.e. simultaneously high longer α -olefin content, high M_{ν} , narrow MWD and high CDBI, impart a number of unique and, in some cases, rather surprising physical, rheological and other properties to the copolymers. As a consequence, the copolymers have a wide number of uses.

FILMS

For structural film applications, the copolymers are generally semicrystalline, having an α -olefin comonomer content below about 12 or 13 mole percent and a density from about 0.88 to about 0.93 g/cm³. The copolymers are formed into film by blown film or extrusion casting procedures using techniques and equipment well known in the linear, low density polyethylene (LLDPE) arts. The present films have high strength and a Young's modulus similar to conventional LLDPE, but have exceptionally high elongation and strain-to-break, and excellent processability due to rheological properties. The films are unusually soft owing to a relatively low storage modulus compared to copolymers made using smaller α -olefin.

The copolymer can be used in a monolayer film, e.g., a film comprised of a single layer of the copolymer without adjacent layers made of a different polymer. Alternatively, the copolymer can be used as one or more layers in a multi-

10

15

20

layer film, e.g. as a structural and/or skin layer. As another option, the copolymer can be used as a cling additive in the cling layer of a film, particularly the amorphous copolymer mentioned above.

include one or more conventional film can additives, e.g. anticling (slip and/or antiblock) additives which may be added during the production of the copolymer or subsequently blended in. Such additives are well-known in the art and include, for example, silicas, silicates, diatomaceous earths, talcs and various lubricants. additives are preferably utilized in amounts ranging from about 100 ppm to about 20,000 ppm, more preferably between about 500 ppm to about 10,000 ppm by weight based upon the weight of the copolymer. The copolymer can, if desired, also include one or more other well-known additives such as, tackifiers. antioxidants, ultraviolet example. absorbers, antistatic agents, release agents, pigments, colorants or the like; however, this again should not be considered a limitation of the present invention.

The film is produced from the ethylene copolymer by any one of a number of well-known extrusion or coextrusion techniques. As preferred examples, any of the blown or chill roll cast processes known in the art can be used.

As previously mentioned, the semicrystalline films of
the present invention have properties making them especially
well suited for use in a variety of applications. For
example, these films can be used in stretch/cling films or
made into other forms, such as a tape, by any one of a
number of well-known cutting, slitting and/or rewinding
operations. Physical properties including, but not limited
to, tensile strength, tear strength and elongation can be
adjusted over wide ranges by altering the copolymer
properties and specifications, as well as additive packages,
as appropriate to meet the requirements to a given wrapping,
bundling, taping or other application.

For bundling, packaging and unitizing applications, the thermoplastic film of the present invention is stretch-wrapped by any one of a number of well-known procedures around an article or a plurality of articles. Typical of

articles suitable for bundling, packaging and unitizing with the present thermoplastic film include, but are not limited to, various foodstuffs (canned or fresh), rolls of carpet, liquid containers and various like goods normally containerized and/or palletized for shipping, storage and/or display.

ADHESIVES

adhesive applications, the polymers of this For invention can be blended with tackifiers and other additives into an adhesive formulation. Suitable tackifiers include those resins which are compatible with the copolymer or Tackifiers are chosen impart copolymer blend. substantial adhesive strength, promote substrate wetting and generally enhance coating performance.

Tackifier components suitable for use in this invention 15 include aliphatic and aromatic hydrocarbon resins such as ESCOREZ or WINGTACK 95. WINGTACK 95 is the tradename for a diene-olefin copolymer of piperylene and 2-methyl-2-butene having a softening point of 95°C. The resin is prepared by the cationic polymerization of 60 weight percent piperylene, 20 10 isoprene, 5 weight percent weight percent cyclopentadiene, 15 weight percent 2-methylbutene and about 10 weight percent dimer. See U. S. Patent 3,577,398. Other tackifying resins of the same general type may be employed in which the resinous copolymer comprises 20-80 weight 25 percent of piperylene and 80-20 weight percent of 2-methyl-Other adhesion-promoting resins which are also in the compositions of this invention include useful hydrogenated rosins, rosin esters, polyterpenes, polymerized and mixed olefins. resins, 30 terpenephenol Hydrogenated hydrocarbon resins obtained under the trade designation ESCOREZ 5380 and ECR-143H are preferred. tackifiers typically have a ring and ball softening point from about 10°C to about 180°C, preferably from about 15°C to about 75°C. Other hydrocarbon tackifiers obtained from Exxon Chemical Co. under the trade designations ECR-111, and ECR-327 have also been found to be particularly preferred. ECR-143H resin, for example, is prepared by the cationic

WO 93/12151 PCT/US92/10590

16

polymerization of a C_5 olefin/diolefin feed stream as described in U. S. Patent 4,916,192 which is hereby incorporated by reference herein.

Pressure sensitive adhesive (PSA) properties are dependent on selection of tackifier resin. Particularly important is the T_g of the tackifier. Tack-related properties can be improved by optimizing the T_g of the PSA system. Selection of tackifier is an important variable in this regard. For example, when tackifiers are blended together, several tack properties can be improved in PSA systems incorporating the blended tackifier over PSA systems incorporating each individual tackifier resin. General tackifier composition is also a strong variable in PSA property optimization.

Adhesive systems which are an embodiment of this invention may contain a tackifier resin in an amount of from about 5 to about 95 parts by weight and the copolymer or copolymers in an amount of from about 5 to about 95 parts by weight, relative to the tackifier parts by weight. Preferred adhesive systems contain the tackifier in an amount of from about 30 to about 70 parts by weight, and copolymer or copolymers in an amount of from about 30 to about 70 parts by weight.

15

20

The adhesive composition may further contain relatively
minor amounts of ingredients such as, oils, fillers,
coupling agents, colorants, antioxidants, and other
stabilizing additives which do not substantially adversely
affect the system such as, for example, by adversely
interfering with adhesion to a substrate surface. The
formulation is preferably a hot-melt essentially free of
solvents and other vaporizable constituents which detract
from the hot-melt characteristics of the formulation, e.g.,
no need for drying or solvent removal.

Coating systems which are an embodiment of this invention may optionally contain a resin including both tackifiers and other high polymers blended up to about 50 parts by weight and the longer-α-olefin/ethylene copolymer in an amount of 50 parts by weight or more wherein the parts by weight of the resin and copolymer components total 100.

10

25

30

35

Antioxidants or stabilizers, when used, can be added at from about 0.1 to about 3 percent by weight, preferably from about 0.1 to about 1.5 percent by weight, more preferably from about 0.1 to about 1 percent by weight, and typically at about 0.5 weight percent.

The optional oils which may be mentioned include refined hydrocarbon oils typically present in adhesives, paraffinic, aromatic, naphthenic and including available under the trade designations KAYDOL (produced by WITCO), TUFFLO (produced by ARCO), and the like. refined oils serve to reduce viscosity and improve surface tack properties.

Particulated fillers which may be also used for thickening and price reduction include glass, amorphous SiO2, fumed alumina, calcium carbonate, fibers and Suitable commercially available fillers are the like. available under the trade designations CAB-O-SIL, ZEOSIL 35, AEROSIL R972, DUCRAL 10 and the like.

Suitable coupling agents include (but are not limited to) organometallic compounds such as, for example, silane-20 organotitanates, organozirconates, compounds, based organozircoaluminates, chrome complexes and the like. These are generally selected to promote adhesion based on the substrates and/or fillers involved in the particular application.

Suitable dyes include Fuchsine (CI 42510), Calcocid Green S (CI 44090), Solvent Yellow 34 (CI 4100B), and the like. Suitable pigments include titanium dioxide, colloidal carbon, graphite, ceramics, clays, phosphor particles and metal particles, e.g. aluminum magnetic iron, copper, and the like.

coating compositions of this invention The preferably prepared as organic solvent solutions of the copolymer and any other components, although copolymer emulsions and hot melts may also be used if so desired. coating compositions may be applied to the substrate from a solution of up to about 40 percent weight solids of the ingredients in a solvent such as toluene, the solvent being removed by evaporation to leave a coating on the substrate

15

20

25

30

surface. Alternatively, the ingredients may be mixed in a solvent, the mixture may be emulsified and the solvent evaporated, and the coating may be applied to a substrate as 50-60 percent weight solids emulsion, the water being removed by evaporation with conventional drying equipment and techniques.

For hot melt application, the coating compositions may be prepared by blending the copolymer with any optional component in the melt until a homogeneous blend is obtained. Various methods of blending materials of this type are known to the art, and any method that produces a homogeneous blend is satisfactory. Typical blending equipment includes, for example, mixing extruders, roll mills, Banbury mixers, Brabenders and the like. In general, the blend components blend easily in the melt and a heated vessel equipped with a stirrer is all that is required. The components are added in no particular order, but generally the copolymer is added first and heated in the vessel until molten. Thereafter, any optional components are then added.

The hot melt formulation may be cooled and later reheated for use, or used directly, e.g. supplied from a reservoir or melt pot to a substrate using conventional equipment, for example, for pumping or pressure extrusion Generally, the hot melt is heated through slot dies. sufficiently for a target viscosity of about 100,000 cps, although a viscosity as high as 150,000 cps can usually be tolerated. For suitable pot stability, the viscosity of the hot melt should not increase more than 20 percent when maintained at the pot temperature for a period of 8 hours. An unusual property of the present copolymers and adhesives formulated therewith is a shear-thinning phenomenon. shear rates at typical hot melt application temperatures, the copolymer has a relatively high viscosity; but at high rates of shear, the viscosity generally declines, usually in This permits the copolymer adhesive to a dramatic fashion. be sprayed onto a substrate surface, e.g. through a nozzle. The copolymer experiences a high shear rate as it passes through the spraying device, typically including relatively small orifices and/or passageways, and requires

15

20

25

30

pressure than a conventional polyolefin, if the conventional polyolefin could be sprayed at all. However, once deposited on the substrate surface where there is very little shear, the viscosity is effectively high and advantageously inhibits running or dripping before the copolymer can cool and solidify.

The preparation of coated articles such as films, sheets, plates and molded objects involves the initial step of coating at least a portion of a surface of the selected article with a solution, emulsion or hot melt of the copolymer or adhesive composition. Any suitable coating technique may be employed while applicable substrates, including composites thereof, may be comprised of paper and paperboard; fiberglass; wood; graphite; conductive metals, e.g. copper, aluminum, zinc, and steel, etc.; and semiconductive substrates such as silicon and gallium arsenide; glass and ceramic; textiles, both natural and synthetic, woven and non-woven; synthetic resins including the homoand copolymers of ethylene, propylene, vinyl chloride, vinylidene chloride, vinyl acetate, styrene, isobutylene, and acrylonitrile; polyvinyl acetal; polyethylene terephthalate; polyamides; and cellulose esters such as cellulose acetate and cellulose butyrate. The latter polymeric substrates may contain fillers or reinforcing agents, such as the various synthetic, natural or modified including, for example, cellulosic fiber, cotton, cellulose acetate, viscose rayon, and paper; glass; and polyamide fibers. These reinforced substrates may be used in laminated or composite form.

The coating of the copolymer or adhesive composition should be applied to the substrate surface so that upon drying its thickness will be in the range of about 0.05 to about 10 mils. Drying of the wet polymer coating may be achieved by air drying or by the application of any other particular drying technique is favored by the practitioner.

A preferred use of the present invention is in the preparation of pressure-sensitive adhesive tapes or in the manufacture of labels. The pressure-sensitive adhesive tape comprises a flexible backing sheet and a layer of the

20

25

35

copolymer or adhesive composition of the novel PSA compound coated on one major surface of the backing sheet. backing sheet may be a plastic film, paper or any other suitable material and the tape may include various other layers or coatings, such as primers, release coatings and the like, which are used in the manufacture of pressuresensitive tapes.

The present coating composition may be used as a corrosion resistant barrier coating on various surfaces in intimate contact with corrosion-causing fluids or gases including water, seawater, high and low pH fluids, and the like or exposed to a corrosion-causing environment. Examples include, liners in food and beverage containers; liners in vessels, pipes, and miscellaneous equipment used in manufacturing plants, ships, and the like; and anti-rust 15 As other useful coatings, coatings for automobiles, etc. the copolymers may be used as film-forming binders or adhesives in the production of various coating and/or impregnating compositions for application to papers and textiles.

OTHER USES

The copolymer of the present invention has a wide number of uses because of its unique properties which can be The copolymer can varied to suit particular applications. film and adhesive in example, for utility, applications, as previously mentioned; in applications requiring super tough polymers with the unique morphology of the present copolymer; in polymer blends as a compatibilizer between normally incompatible polymers; in film surface modifications wherein the copolymer is added to or coated on, e.g. a conventional polyethylene, and the film surface can also be subjected to corona discharge or other surface treatment; in polymer processing as an additive to enhance the melt viscosity of the thermoplastic, elastomer thermoplastic elastomer being processed; in soft elastomer applications, particularly vulcanizable elastomers wherein imparts termonomer which copolymer includes a the tactile requiring applications vulcanizability; in

25

3.0

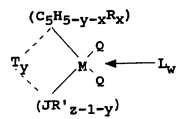
•

polymer; in various molding applications, e.g. injection molding, blow molding and thermoforming; and the like.

CATALYST COMPONENT

The present invention relates to copolymers of ethylene lpha-olefins made by a process comprising longer 5 polymerizing the longer α -olefins with ethylene in the presence of a catalyst providing a low ethylene:comonomer reactivity ratio, preferably a ratio less than about 50, more preferably less than about 30, especially from about 3 to about 20, and more particularly from about 5 to about 15. activated an comprises catalyst preferred cyclopentadienyl-transition metal compound wherein transition metal component is from Group IV B.

The Group IV B transition metal component of the catalyst system is represented by the general formula: 15



wherein: M is Zr, Hf or Ti and is in its highest formal 20 oxidation state (+4, d0 complex);

 $(C_5H_{5-y-x}.R_x)$ is a cyclopentadienyl ring which substituted with from zero to five substituent groups R, "x" is 0, 1, 2, 3, 4 or 5 denoting the degree of substitution, and each substituent group R is, independently, a radical selected from a group consisting of C_1-C_{20} hydrocarbyl radicals, substituted C_1 - C_{20} hydrocarbyl radicals wherein one or more hydrogen atoms is replaced by a halogen radical, an amido radical, a phosphido radical, an alkoxyl radical or any other radical containing a Lewis acidic or basic hydrocarbyl-substituted metalloid C1-C20 functionality, radicals wherein the metalloid is selected from the Group IV A of the Periodic Table of Elements, and halogen radicals, amido radicals, phosphido radicals, alkoxy radicals,

15

20

25

30

alkylborido radicals or any other radical containing a Lewis acidic or basic functionality or $(C_5H_{5-\gamma-x}R_x)$ is a cyclopentadienyl ring in which two adjacent R-groups are joined forming C_4-C_{20} ring to give a saturated or unsaturated polycyclic cyclopentadienyl ligand such as indenyl, tetrahydroindenyl, fluorenyl or octahydrofluorenyl;

(JR' $_{z-1-y}$) is a heteroatom ligand in which J is an element with a coordination number of three from Group V A or an element with a coordination number of two from Group VI A of the Periodic Table of Elements, preferably nitrogen, phosphorus, oxygen or sulfur with nitrogen being preferred, and each R' is, independently a radical selected from a group consisting of C_1 - C_{20} hydrocarbyl radicals, substituted C_1 - C_{20} hydrocarbyl radicals wherein one or more hydrogen atoms is replaced by a halogen radical, an amido radical, a phosphido radical, an alkoxy radical or any other radical containing a Lewis acidic or basic functionality, and "z" is the coordination number of the element J;

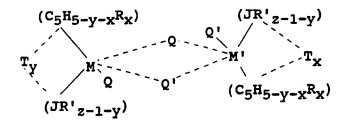
Each Q is, independently any univalent anionic ligand such as halogen, hydride, or substituted or unsubstituted C_1 - C_{20} hydrocarbyl, alkoxide, aryloxide, amide, arylamide, phosphide or arylphosphide, provided that where any Q is a hydrocarbyl such Q is different from $(C_5H_{5-\gamma-x}R_x)$ or both Q together may be an alkylidene or a cyclometallated hydrocarbyl or any other divalent anionic chelating ligand.

"y" is 0 or 1 when w is greater than 0; y is 1 when w is 0; when "y" is 1, T is a covalent bridging group containing a Group IV A or V A element such as, but not limited to, a dialkyl, alkylaryl or diaryl silicon or germanium radical, alkyl or aryl phosphine or amine radical, or a hydrocarbyl radical such as methylene, ethylene and the like.

L is a Lewis base such as diethylether, tetraethylammonium chloride, tetrahydrofuran, dimethylaniline, aniline, trimethylphosphine, n-butylamine, and the like; and "w" is a number from 0 to 3; L can also be a second transition metal compound of the same type such that the two metal centers M and M' are bridged by Q and Q', wherein M' has the same meaning as M and Q' has the same

10

meaning as Q. Such compounds are represented by the formula:



A preferred activator is an alumoxane component which may be represented by the formulas: $(R^3-Al-O)_m$; $R^4(R^5-Al-O)_m$ -AlR 6_2 or mixtures thereof, wherein R^3-R^6 are, independently, a univalent anionic ligand such as a C_1-C_5 alkyl group or halide and "m" is an integer ranging from 1 to about 50 and preferably is from about 13 to about 25.

Examples of the T group which are suitable as a constituent group of the Group IV B transition metal component of the catalyst system are identified in Column 1 of Table 1 under the heading "T".

(when y=1)				
	(C5H _{F-V-X} R _V)	(18,-1)	o	-
	cyclopentadienyl	t-butvlemido	ŀ	E
	methylcyclopentadienyl	shows omide	nyaride	zirconium
di-n-propylsilyl	1.2-dimethylevelopentadiamyl	Men y Callingo	chloro	hafnium
ditsopropylativi	1 Z - 42 - 44 - 44 - 44 - 44 - 44 - 44 -	p-n-putylphenylamido	methyl	titanium
di-n-butyisilvi	1,3-unitectifyteyetopentaatenyt	cyclohexylamido	ethyl	
4	Indenyt	Perflurophenylamido	phenyl	
V	1,2-diethylcyclopentadienyl	n-butytamido	fluoro	
dı- <u>n</u> -hexylaflyl	tetramethylcyclopentadienyl	methylamido	bromo	
methylphenylsilyl	ethylcyclopentadienyl	ethylemido	ioda	
ethylmethylsilyl	n-butyl cyclopentadienyl	n-propytamido	n-pronvl	
diphenylsilyl	cyclohextmethylcyclopentadienyl	isopropylamido	i connonsi	
di(p-t-butylphenethylsilyl)	n-octyl cyclopentadienyl	benzvlamido	Taylor Day	
<u>n</u> -hexylmethylsilyl	beta-phenyl propyl cyclopentadienyl	t-butylphosphido	n-twick t	
cyclopentamethylenesilyl	tetrahydroindenyl	ethylphosphido	7 Line	
cyclotetramethylenesilyl	propylcyclopentadienyl	ohenvl phosphido	1 SOBIITY L	
cyclotrimethylenesilyl	t-butylcyclopetnadienyl	cyclohexylphosphido	- topator	
dimethylgermanyl	benzylcyclopentadienyl	oxo (when v = 1)	hentul	
diethylgermanyl	diphenylmethylcyclopentadienyl	sulfide (upen v = 1)	Tebra Control	
	trimethylgermylcyclopentadienyl	methoxide (when v = 0)	non/	
	trimethylstamylcyclopentadienyl	ethoxide (when v = 0)	- Control	
	triethylplumbylcyclopentadienyl	methylthio (when y = 0)	Getyl	
t-butylphosphido	trifluromethylcyclopentadienyl	ethylthio (when y = 0)	methoxy	
ethytphosphido	trimethylsilylcyclopentadienyl		ethox.	
phenylphosphido	pentamethylcyclopentadienyl (when y = 0)		Propositi	
			Putoni	
dimethylene	octahydrofluorenyl		phenous	
diethylmethylene	N.Mdimethylamidocyclopentadienyl		dimethyl parids	
	dimethylphosphidocyclopentadienyl		diethol emido	
dimethylethylene	methoxycyclopentadienyl		methyl athyl omide	

TABLE 1

TABLE 1 (CONT'D)

T (when y≃1)	(C5H5-y-xRx)	(JR'z-1-y)	σ	Σ
dipropylethylene	(N, N-dimethylamidomethyl)cyclopentadienyl		diphenylamido	
propylene			diphenylphosphido	
dimethylpropylene			dicyclohexylphosphido	
diethylpropylene			dimethytphosphido	
1,1-dimethyl-3,3-			methylidene (both Q)	
dimethylpropylene				
tetramethyldisiloxane			ethylidene (both 0)	
1,1,4,4-			propylidene (both q)	
tetramethyldisilylethylene				
			ethyleneglycoldianion	
			(both 0)	

15

20

25

30

Exemplary hydrocarbyl radicals for the Q are methyl, ethyl, propyl, butyl, amyl, isoamyl, hexyl, isobutyl, heptyl, octyl, nonyl, decyl, cetyl, 2-ethylhexyl, phenyl and the like, with methyl being preferred. Exemplary halogen atoms for Q include chlorine, bromine, fluorine, and iodine, with chlorine being preferred. Exemplary alkoxides and aryloxides for Q are methoxide, phenoxide and substituted phenoxides such as 4-methylphenoxide. Exemplary amides for O are dimethylamide, diethylamide, methylethylamide, di-tbutylamide, diiospropylamide and the like. Exemplary amides for Q are dimethylamide, diethylamide, methylethylamide, dit-butylamide, diisopropylamide and the like. Exemplary aryl amides are diphenylamide and any other substituted phenyl Exemplary phosphides for Q are diphenylphosphide, dicyclohexylphosphide, diethylphosphide, dimethylphosphide Exemplary alkyldiene radicals for both Q and the like. are methylidene, ethylidene and propylidene. together Examples of the Q group which are suitable as a constituent group or element of the Group IV B transition metalcomponent of the catalyst system are identified in Column 4 of Table 1 under the heading "Q".

Suitable hydrocarbyl and substituted hydrocarbyl radicals, which may be substituted as an R group for at least one hydrogen atom in the cyclopentadienyl ring, will contain from 1 to about 20 carbon atoms and include straight and branched alkyl radicals, cyclic hydrocarbon radicals, alkyl-substituted cyclic hydrocarbon radicals, aromatic radicals, alkyl-substituted aromatic radicals, phosphido hydrocarbon radicals, alkoxy substituted substituted hydrocarbon radicals, alkylborido substituted radicals and cyclopentadienyl rings containing one or more Suitable organometallic saturated or unsaturated rings. radicals, which may be substituted as an R group for at least one hydrogen atom in the cyclopentadienyl ring, include trimethylsilyl, triethylsilyl, ethyldimethylsilyl, methyldiethylsilyl, triphenylgermyl, trimethylgermyl and the like. Other suitable radicals that may be substituted for one or more hydrogen atom in the cyclopentadienyl ring halogen radicals, amido radicals, phosphido include

15

radicals, alkoxy radicals, alkylborido radicals and the like. Examples of cyclopentadienyl ring groups $(C_5H_5._{y-x}R_x)$ which are suitable as a constituent group of the Group IV B transition metal component of the catalyst system are identified in Column 2 of Table 1 under the heading $(C_5H_5._{y-x}R_x)$.

Suitable hydrocarbyl and substituted hydrocarbyl radicals, which may be used as an R' group in the heteroatom J ligand group, will contain from 1 to about 20 carbon atoms and include straight and branched alkyl radicals, cyclic hydrocarbon radicals, alkyl-substituted cyclic hydrocarbon radicals, aromatic radicals, alkyl-substituted aromatic radicals, halogen radicals, amido radicals, phosphido radicals, alkylborido radicals and the like. Examples of heteroatom ligand groups (JR'2-1-v) which are suitable as a constituent group of the Group IV B transition metal component of the catalyst system re identified in Column 3 of Table 1 under the heading (JR'2-1-v).

Table 1 depicts representative constituent moieties for -20 the "Group IV B transition metal component", the list is for illustrative purposes only and should not be construed to be limiting in any way. A number of final components may be formed by permuting all possible combinations constituent moieties with each other. Illustrative 25 compounds are: dimethylsilyltetramethyl-cyclopentadienyltert-butylamido zirconium dichloride. dimethylsilytetramethylcyclopentadienyl-tert-butylamido hafnium dichloride, dimethylsilyl-<u>tert</u>-butylcyclopentadienyl-tert-butylamido hafnium dichloride. 30 dimethylsilyltrimethylsilylcyclopentadienyl-tert-butylamido zirconium dichloride, dimethylsilyltetramethylcyclopentadienylphenylamido zirconium dichloride, dimethylsilyltetramethylcyclopentadienyl-phenylamido hafnium dichloride, methylphenylsilyl-tetramethylcyclopentadienyl-35 tert-butylamido zirconium dichloride, methylphenylsilyltetramethylcyclopentadienyl-tert-butylamido hafnium dichloride, methylphenylsilyltetramethylcyclopentadienyl-tert-butylamido hafnium dimethylsilyltetramethylcyclopentadienyl-p-ndimethyl,

25

30

35

dimethylsilyldichloride, zirconium butylphenylamido ${\tt tetramethylcyclopentadienyl-p-\underline{n}-butylphenylamido}$ hafnium dichloride.

As noted, titanium species of the Group IV B transition metal compound have generally been found to yield catalyst systems which in comparison to their zirconium or hafnium analogues, are of higher activity and α -olefin comonomer Illustrative, but not limiting of incorporating ability. the titanium species which exhibit such superior properties methylphenylsilyltetramethylcyclopentadienyl-tert-10 are dichloride, dimethylsilylbutylamido titanium tetramethylcyclopentadienyl-p-n-butylphenylamido titanium dimethylsilyltetramethylcyclopentadienyl-pdichloride, methoxyphenylamido titanium dichloride, dimethylsilyl-tertbutylcyclopentadienyl-2,5-di-tert-butylphenylamido titanium dimethylsilylindenyl-tert-butylamido titanium dichloride, dimethylsilyltetramethyldichloride, dichloride, cyclopentadienylcyclohexylamido titanium dimethylsilylfluorenylcyclohexylamido titanium dichloride, dimethylsilyltetramethylcyclopentadienyl-phenylamido dimethylsilyldichloride, titanium tetramethylcyclopentadienyl-<u>tert</u>-butylamido titanium dimethylsilyltetramethylcyclopentadienyldichloride, cyclododecylamido titanium dichloride, and the like.

For illustrative purposes, the above compounds and those permuted from Table 1 do not include the Lewis base ligand (L). The conditions under which complexes containing Lewis base ligands such as ether or those which form dimers is determined by the steric bulk of the ligands about the t-butyl group For example, the center. metal $Me_2Si(Me_4C_5)(N-\underline{t}-Bu)ZrCl_2$ has greater steric requirements than the phenyl group in Me₂Si(Me₄C₅)(NPh)ZrCl₂.Et₂O thereby not permitting ether coordination in the former compound. due to the decreased steric bulk of Similarly, trimethylsilylcyclopentadienyl group in [$Me_2Si(Me_3SiC_5H_3)(N-m)$] <u>t</u>-Bu) ZrCl₂]₂ versus that of the tetramethylcyclopentadienyl group in $Me_2Si(Me_4C_5)(N-\underline{t}-Bu)ZrCl_2$, the former compound is dimeric and the latter is not.

15

20

25

30

35

Generally the bridged species of the Group transition metal compound ("y"=1) are preferred. compounds can be prepared by reacting a cyclopentadienyl lithium compound with a dihalo compound whereupon a lithium halide salt is liberated and a monohalo substituent is covalently bound to the cyclopentadienyl compound. The substituted cyclopentadienyl reaction product reacted with a lithium salt of a phosphide, oxide, sulfide or amide (for the sake of illustrative purposes, a lithium halo element of the monohalo whereupon the amide) substituent group of the reaction product reacts to liberate a lithium halide salt and the amine moiety of the lithium amide salt is covalently bound to the substituent of the cyclopentadienyl reaction product. The resulting amine derivative of the cyclopentadienyl product is then reacted with an alkyl lithium reagent whereupon the labile hydrogen atoms, at the carbon atom of the cyclopentadienyl compound and at the nitrogen atom of the amine moiety covalently bound to the substituent group, react with the alkyl of the lithium alkyl reagent to liberate the alkane and produce a dilithium salt of the cyclopentadienyl compound. the bridged species of the Group IV B transition metal compound is produced by reacting the dilithium cyclopentadienyl compound with a Group IV B transition metal preferably a Group IV B transition metal halide.

Unbridged species of the Group IV B transition metal compound can be prepared from the reaction of a cyclopentadienyl lithium compound and a lithium salt of an amine with a Group IV B transition metal halide.

Suitable, but not limiting, Group IV B transition metal compounds which may be utilized in the catalyst system of this invention include those bridged species ("y"=1) wherein the T group bridge is a dialkyl, diaryl or alkylaryl silane, or methylene or ethylene. Exemplary of the more preferred species of bridged Group IV B transition metal compounds are dimethylsilyl, methylphenylsilyl, diethylsilyl, ethylphenylsilyl, diphenylsilyl, ethylene or methylene bridged compounds. Most preferred of the bridged species

20

25

30

are dimethylsilyl, diethylsilyl and methylphenylsilyl bridged compounds.

Suitable Group IV B transition metal compounds which are illustrative of the unbridged ("y"=0) species which may be utilized in the catalyst systems of this invention are exemplified by pentamethylcyclopentadienyldi-t-butylphosphinodimethyl hafnium; pentamethylcyclopentadienyldi-t-butylphosphinomethylethyl hafnium; cyclopentadienyl-2-methylbutoxide dimethyl titanium.

To illustrate members of the Group IV B transition metal component, select any combination of the species in Table 1. An example of a bridged species would be dimethylsilylcyclopentadienyl- \underline{t} -butylamidodichloro

15 zirconium; an example of an unbridged species would be cyclopentadienyldi- \underline{t} -butylamidodichloro zirconium.

Those species of the Group IV B transition metal component wherein the metal is titanium have been found to impart beneficial properties to a catalyst system which are unexpected in view of what is known about the properties of which are compounds bis(cyclopentadienyl) titanium Whereas titanocenes in their cocatalyzed by alumoxanes. soluble form are generally unstable in the presence of aluminum alkyls, the monocyclopentadienyl titanium metal components of this invention, particularly those wherein the heteroatom is nitrogen, generally exhibit greater stability presence of aluminum alkyls and higher catalyst in the activity rates.

Further, the titanium species of the Group IV B transition metal component catalyst of this invention generally exhibit higher catalyst activities and the production of polymers of greater molecular weight than catalyst systems prepared with the zirconium or hafnium species of the Group IV B transition metal component.

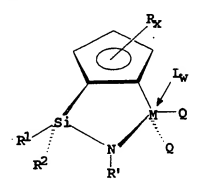
Generally, wherein it is desired to produce an α -olefin copolymer which incorporates a high content of α -olefin, while maintaining high molecular weight polymer the species of Group IV B transition metal compound preferred is one of

20

25

30

The most preferred species of titanium metal titanium. compounds are represented by the formula:



wherein Q, L, R', R, "x" and "w" are as previously defined and \mathbb{R}^1 and \mathbb{R}^2 are each independently a C_1 to C_{20} hydrocarbyl radicals, substituted C1 and C20 hydrocarbyl radicals wherein one or more hydrogen atom is replaced by a halogen atom; R2 and \mathbb{R}^3 may also be joined forming a C_3 to C_{20} ring which incorporates the silicon bridged. Suitable hydrocarbyl and substituted hydrocarbyl radicals which may be used as an R' group have been described previously. Preferred R' groups include those bearing primary carbons bonded directly to the nitrogen atom such as methyl, ethyl, n-propyl, n-butyl, nn-decyl, n-dodecyl, n-tetradecyl, 15 hexyl, n-octyl, hexadecyl, n-octadecyl, benzyl and the like, and those bearing secondary carbons bonded directly to the nitrogen atom such as 2-propyl, 2-butyl, 3-pentyl, 2-heptyl, 2-octyl, cyclobutyl, cyclopentyl, cyclohexyl, cyclopropyl, cycloheptyl, cyclooctyl, cyclododecyl, 2-norbornyl and the like.

Also, the most preferred cyclopentadienyl ring is tetramethylcyclopentadiene (R = Me and x = 4).

The alumoxane component of the catalyst system is an oligomeric compound which may be represented by the general formula $(R^3-Al-0)_m$ which is a cyclic compound, or may be $R^4(R^5-Al-O-)_m-AlR^6_2$ which is a linear compound. An alumoxane is generally a mixture of both the linear and cyclic compounds. In the general alumoxane formula R^3 , R^4 , R^5 , and R6 are, independently a univalent anionic ligand such as a C_1-C_5 alkyl radical, for example, methyl, ethyl, propyl, butyl, pentyl or halide and "m" is an integer from 1 to abut

10

30

35

50. Most preferably, R^3 , R^4 , R^5 and R^6 are each methyl and "m" is at least 4. When an alkyl aluminum halide is employed in the preparation of alumoxane, one or more of R^{3-6} could be halide.

As is now well known, alumoxanes can be prepared by various procedures. For example, a trialkyl aluminum may be reacted with water, in the form of a moist inert organic solvent; or the trialkyl aluminum may be contacted with a hydrated salt, such as hydrated copper sulfate suspended in an inert organic solvent, to yield an alumoxane. Generally, however prepared, the reaction of a trialkyl aluminum with a limited amount of water yields a mixture of both the linear and cyclic species of alumoxane.

Suitable alumoxanes which may be utilized in the catalyst systems of this invention are those prepared by the 15 alkylaluminum reagent; such a hydrolysis of trimethylaluminum, triethyaluminum, tripropylaluminum, dimethylaluminumchloride, triisobutylaluminum, diisobutylaluminumchloride, diethylaluminumchloride, and the Mixtures of different alkyl aluminum reagents in 20 preparing an alumoxane may also be used. The most preferred alumoxane for use is methylalumoxane (MAO), particularly methylalumoxanes having a reported average degree oligomerization of from about 4 to about 25 ("m"=4 to 25) with a range of 13 to 25 being most preferred. 25

As an alternative to the alumoxane activation, the metallocene component can be ionically activated using the procedures and techniques set forth in Turner et al., U. S. Ser. No. 133,052, filed December 21, 1987; Turner et al., U. S. Ser. No. 133,480, filed December 22, 1987; Greg et al., U. S. Ser. No. 542,236, filed June 22, 1990; and EP Publication Nos. 277,004; 418,044; and 426,637; all of which are hereby incorporated by reference. Briefly, for ionic activation, the metallocene has at least one substituent capable of reacting with a proton. The metallocene is activated by reaction with a proton-donating cation and a bulky, non-coordinating anion which stabilizes the metal reaction. cation formed by the metallocene-proton Typically, Q in the above formula is hydrocarbyl, the cation

15

is trialkylammonium, for example, and the anion is tetraperfluorophenyl borate, for example.

CATALYST SYSTEMS

The catalyst systems employed in the method of the invention comprise a complex formed upon admixture of the Group IV B transition metal component with an activating component. The catalyst system may be prepared by addition of the requisite Group IV B transition metal and alumoxane components, or a previously cationically activated Group IV B transition metal component, to an inert solvent in which olefin polymerization can be carried out by a solution, slurry or bulk phase polymerization procedure.

The catalyst system may be conveniently prepared by placing the selected Group IV B transition metal component and the selected alumoxane or ionic activating component(s), in an alkane or aromatic any order of addition, hydrocarbon solvent, preferably one which is also suitable service as a polymerization diluent. Where the hydrocarbon solvent utilized is also suitable for use as a polymerization diluent, the catalyst system may be prepared in situ in the polymerization reactor. Alternatively, the catalyst system may be separately prepared, in concentrated form, and added to the polymerization diluent in a reactor. Or, if desired, the components of the catalyst system may be and added solutions separate prepared as polymerization diluent in a reactor, in appropriate ratios, as is suitable for a continuous liquid polymerization Alkane and aromatic hydrocarbons reaction procedure. suitable as solvents for formation of the catalyst system and also as a polymerization diluent are exemplified by, but are not necessarily limited to, straight and branched chain hydrocarbons such as isobutane, butane, pentane, hexane, cyclic like, and alicyclic and the octane heptane, cyclohexane, cycloheptane, such as hydrocarbons methylcycloheptane and the like, and methylcyclohexane, aromatic and alkyl-substituted aromatic compounds such as benzene, toluene, xylene and the like. Suitable solvents also include liquid olefins which may act as monomers or

15

20

25

30

comonomers, including ethylene, propylene, 1-butene, 1-hexene and the like, particularly when the catalyst components are prepared as separate solutions.

In accordance with this invention optimum results are generally obtained wherein the Group IV B transition metal compound is present in the polymerization diluent in a concentration of preferably from abut 0.00001 to about 10.0 millimoles/liter of diluent and the alumoxane component, when used, is present in an amount to provide a molar aluminum to transition metal ratio of from about 0.5:1 to about 20,000:1. Sufficient solvent is normally used so as to provide adequate heat transfer away from the catalyst components during reaction and to permit good mixing.

The catalyst system ingredients, that is, the Group IV B transition metal, the alumoxane and/or ionic activators, and polymerization diluent, can be added to the reaction vessel rapidly or slowly. The temperature maintained during the contact of the catalyst components can vary widely, such as, for example, from -100°C to 300°C. Greater or lesser temperatures can also be employed. Preferably, during formation of the catalyst system, the reaction is maintained within a temperature of from about 25°C to 100°C, most preferably about 25°C.

individual catalyst the times, At all components, as well as the catalyst system once formed, are protected from oxygen and moisture. Therefore, reactions are performed in an oxygen and moisture free atmosphere and, where the catalyst system is recovered separately it is recovered in an oxygen and moisture free Preferably, therefore, the reactions atmosphere. performed in the presence of an inert dry gas such as, for example, helium or nitrogen.

POLYMERIZATION PROCESS

In a preferred embodiment of the process of this invention the catalyst system is utilized in the liquid phase (slurry, solution, suspension or bulk phase or combination thereof), high pressure fluid phase or gas phase polymerization of an olefin monomer. These processes may be

35

employed singularly or in series. The liquid phase process comprises the steps of contacting a longer α -olefin monomer and ethylene with the catalyst system in a suitable polymerization diluent and reacting said monomers in the presence of said catalyst system for a time and at a temperature sufficient to produce a polyolefin of high molecular weight. Conditions most preferred for copolymerization of ethylene are those wherein ethylene is submitted to the reaction zone at pressures of from about 0.019 psia to about 50,000 psia and the reaction temperature 10 is maintained at from about -100°C to about 300°C. aluminum to transition metal molar ratio is preferably from about 1:1 to 18,000 to 1. A more preferable range would be 1:1 to 2000:1. The reaction time is preferably from about 10 seconds to about 4 hours. 15 Without limiting in any way the scope of the invention, one means for carrying out the process of the present invention for production of a copolymer is as follows: in a stirred-tank reactor liquid α -olefin monomer is introduced, such as 1-dodecene. 20 catalyst system is introduced via nozzles in either the vapor or liquid phase. Feed ethylene gas is introduced either into the vapor phase of the reactor, or sparged into the liquid phase as is well known in the art. The reactor contains a liquid phase composed substantially of liquid α -25 olefin comonomer, together with dissolved ethylene gas, and a vapor phase containing vapors of all monomers. The reactor temperature and pressure may be controlled via reflux of vaporizing α -olefin monomer (autorefrigeration), well as by cooling coils, jackets The polymerization rate is generally controlled the concentration of catalyst. The ethylene content of the polymer product is determined by the ratio of ethylene to α olefin comonomer in the reactor, which is controlled by manipulating the relative feed rates of these components to the reactor.

As before noted, a catalyst system wherein the Group IV B transition metal component is a titanium species has the ability to incorporate high contents of longer α -olefin Accordingly, the selection of the Group IV B comonomers.

transition metal component is another parameter which may be utilized as a control over the ethylene content of a copolymer within a reasonable ratio of ethylene to longer α -olefin comonomer.

EXAMPLES

CATALYST PREPARATION

All catalyst preparation and polymerization procedures 5 were performed under an inert atmosphere of helium or nitrogen. Solvent choices were often optional, for example, in most cases either pentane or 30-60 petroleum ether could The choice between tetrahydrofuran (THF) be interchanged. and diethyl ether was a bit more restricted, but in several 10 reactions, either could be used. The lithiated amides were prepared from the corresponding amines and either nmethyllithium (n-BuLi) or butyllithium Tetramethylcyclopentadienyl-lithium (C5Me4HLi) was prepared according to the procedures of C. M. Fendrick et al., 15 Organometallics, 1984, 3, 819 and F. H. Kohler and K. H. Other lithiated Z Naturforsch, 1982, 376, 144. cyclopentadienyl compounds were substituted prepared from the corresponding cyclopentadienyl ligand and n-BuLi or MeLi, or by reaction of MeLi with the proper 20 TiCl4 was typically used in its etherate form. The etherate was generally prepared by simply adding $TiCl_{\lambda}$ to ether, filtering off the solid product and vacuum drying. substituted amines, silanes, ZrCl4, HfCl4, TiCl, unsubstituted cyclopentadienyl compounds or precursors, and lithium reagents were purchased from Aldrich Chemical Methylalumoxane was supplied Company or Petrarch Systems. by either Schering or Ethyl Corporation.

C₅Me₄HLi (10.0 g, 0.078 mol) was slowly added to Me₂SiCl₂ (11.5 ml, 0.095 mol, in 225 ml of THF solution). The solution was stirred for 1 hour to assure a complete reaction. The solvent was then removed in vacuo. Pentane was added to precipitate the LiCl. The mixture was filtered through diatomaceous earth and the solvent was removed from the

15

20

25

Tetramethylcyclopentadienyldimethylchlorosilane, (C₅Me₄H)SiMe₂Cl, (15.34 g, 0.071 mol) was recovered as a pale yellow liquid.

 $(C_5 Me_4 H) SiMe_2 Cl$ (8.0 g, 0.037 mol) was slowly added to a suspension of lithium cyclododecylamine (LiHNC₁₂H₂₃) (7.0 g, 0.037 mol, ~80 ml THF). The mixture was stirred overnight. The THF was then removed by vacuum to a cold trap held at ~196°C. A mixture of petroleum ether and toluene was added to precipitate the LiCl. The mixture was filtered through diatomaceous earth. The solvent was removed from the filtrate.

Tetramethylcyclopentadienyl aminocyclododecyldimethylsilane, $Me_2Si(C_5 Me_4 H) (NHC_{12}H_{23})$, (11.8 g, 0.033 mol) was isolated as a pale yellow liquid.

 $Me_2Si(C_5Me_4H) (NHC_{12}H_{23})$ (11.9 g, 0.033 mol) was diluted with ~150 ml of ether. MeLi (1.4 M, 47 ml, 0.066 mol) was added slowly, and the mixture was stirred for 2 hours. The ether was reduced in volume by evaporation. The product was filtered off. The product [Me_2Si(C_5Me_4)(NC_{12}H_{23})]Li_2, was washed with several small portions of ether, then vacuum dried to yield 11.1 g (0.030 mol).

0.008 mol) (3.0 g, was $[Me_2Si(C_5Me_4)(NC_{12}H_{23})]Li_2$ suspended in cold ether. TiCl₄·2Et,0 (2.7 g, 0.008 mol) was slowly added and the resulting mixture was The ether was removed via a vacuum to a cold overnight. Methylene chloride was added to trap held at -196°C. precipitate the LiCl. The mixture was filtered through diatomaceous earth. The solvent was significantly reduced in volume by evaporation and petroleum ether was added to This mixture was refrigerated precipitate the product. prior to filtration in order to maximize precipitation. solid collected was recrystallized from methylene chloride and $Me_2Si(C_5Me_4)(NC_{12}H_{23})TiCl_2$ was isolated (1.0 g, 2.1 mmol).

POLYMERIZATION EXAMPLES 1-21

Polymerization was done in a 1-liter autoclave reactor equipped with a paddle stirrer, an external water jacket for temperature control, a regulated supply of dry nitrogen, ethylene, propylene, 1-butene and hexane, and a septum inlet for introduction of other solvents or comonomers, transition

metal compound and alumoxane solutions. The reactor was dried and degassed thoroughly prior to use. A typical run consisted of injecting a quantity of solvent (typically toluene), the comonomer and 1.0 M methylalumoxane (MAO) into the reactor. The reactor was then heated to the specified temperature and the transition metal compound solution and the ethylene at a pressure of 4.08 atm were introduced into the system. The polymerization reaction was limited to a specified time, typically 30 minutes. The reaction was ceased by rapidly cooling and venting the system, and the resulting polymer was recovered by evaporating the solvent under a stream of nitrogen.

Remaining process run conditions are given in Table 2 including the amount of transition metal catalyst solution (TMC) used, the amount of methylalumoxane solution used, the Al/Ti molar ratio, the amount of toluene and comonomer used, the polymerization temperature, polymer yield, catalyst efficiencies in terms of kg polymer per mole catalyst atm hr and kg polymer per mole catalyst reactivity ratio.

For example, 390 ml of toluene, 6 ml of 1 M MAO and 10 ml of 1-decene were added to the reactor described above. The reactor was heated to 80°C prior to introducing 1.2 ml of the catalyst stock solution made by dissolving 13.5 mg of the transition metal compound in 10 ml of toluene. reactor was then immediately pressurized with 4.08 atm of The polymerization reaction was limited to 30 minutes after which time the reaction was ceased by rapidly The resulting polymer (39 cooling and venting the system. g) was recovered by evaporating the solvent under a stream 30 of nitrogen. Catalyst productivity was calculated at 5,212 (kg polymer/mol TMC atm hr) and 23,038 (kg polymer/mol Polymer characteristics include a GPC/DRI TMC·hr). molecular weight of 123,000 daltons, a molecular weight distribution of 2.6, 3.2 mole percent incorporated 1-decene giving a catalyst reactivity ratio of 18.7 ethylene to 1decene, a polymer density of 0.914 g/ml, a melting point of 118°C and a T_g of -100°C(T_{α}) and -70°C(T_{β}).

39 TABLE 2

				7112	AIG4	TOLLENE	OLE	POLY-	PRODUC-	PRODUC-
Ex	OLEFIN USED	TMC	TMC	TMC Stock	AVM	(ml)	-FIN	MER	TIVITY(kg	TIVITY
	ļ	Stock	Stock	Used		(11)	(ml)	YIELD	P/mol TMC	(kg P/mol
1		(mg/ 10ml)	(ml)	(mg)				(9)	-atm -hr)	TMC-hr)
2	1-decene	13.5	1	1.35	2127	395	5	20	3475	14,177
1-2	1-devene					200	10	39	5647	23.038
3	1-decene	13.5	1.2	1.62	1772	390	10	35		
4	1-decene	13.5	1	1.35	2127	380	20	40	6950	28,354
5	1-decene	13	11	1.30	2208	350	50	52	9382	38,278
6	1-dodecene	13.4	1	1.34	2142	395	5	10	1750	7,141
7	1-dodecene	13.5	1	1.35	2127	390	10	88	15,289	62,380
В	1-dodecene	13.4	1	1.34	2142	375	25	45	7,877	32,137
9	1-dodecene	13.4	1	1.34	2142	350	50	60	10,502	42,849
10	1-dodecene	13.4	1	1.34	2142	300	100	70	12,253	49,990
11	1-tetradecene	13.5	1	1.35	2127	395	5	32	5560	22,683
12	1-tetradecene	13.5	1	1.35	2127	390	10	41	7123	29,063
13	1-tetradecene	13.4	1	1.34	2142	375	25	35	6126	24,995
14	1-tetradecene	13.4	1	1.34	2142	350	50	40	7001	28,566
15	1-hexadecene	13.5	1	1.35	2127	395	5	17	2954	12,051
16	1-hexadecene	13.5	1	1.35	2127	390	10	22	3822	15,595
17	1-hexadecene	13.5	1	1.35	2127	380	20	30	5212	21,266
18	1-octadecene	13.5	1	1.35	2127	395	5	12	2085	8,506
-		13.5	1	1.35	2127	390	10	16	2780	11,342
19	1-octadecene	13.3			-				4244	17 721
20	1-octadecene	13.5	1	1.35	2127	380	20	25	4344	17,721
21	1-octadecene	12	1	1.20	2392	350	50	47	9187	37,481

Resulting polymer characteristics are given in Table 3 including weight average molecular weight, molecular weight 5 distribution, comonomer concentration, polymer density, melting point and glass transition temperatures (both \textbf{T}_{α} and T_{β}).

40

ma	BLE	2
'I'A	BLE	

				1	'ABLL	٤ .				
EX	OLEFIN USED	MW ³ (daltons)	MWD	mol% a- OLEFIN	rı	POLY- MER DENSITY (g/ml)	Tm (°C)	Tg (°C) (T _g ,T _b)	(psi)	STRAIN TO BREAK (%)
2	1-deceme	85,000	2.3	3.8	7.8	0.929		-106/ -22	7800	590
3	1-decene	123,100	2.6	3.2	18.7	0.914	118	-100/ -70	7850	487
4	1-decene	108,200	2.2	8.2	13.8	0.895	115	<u> </u>	2925	540
5	1-decene	186,400/ 299,500	2.5	-	-			-		
6	1-dodecene	120,000	2.1	3.5*	7.3	0.928	87	-102/-14	3960	100
7	1-dodecene	118,200	4.7	3.7	13.7	0.920	109		10,390	704
8	1-dodecene	112,000	2.2	11.5	10.1	0.889	92	-105/-20	550	539
9	1-dodecene	139,000	2.3	19.5	10.9	<u> </u>	<u> -</u>	- /-30		_
10	1-dodecene	94,000/	2.0	29.4	12.6	<0.86	-23	•		-
11	1-tetradecene	117,000	2.6	1.8	12.6	0.933	124		22,040	614
12	1-tetradecene	121,600	2.7	4.1	10.8	0.924		-	16,490	600
13	1-tetradecene	90,000	2.0	10.4	9.9	0.883	-7	-102/-11		
14	1-tetradecene	73,000/ 140.000	2.0	17.8	10.7	-	4	-		
15	1-hexadecene	88,000	2.1	3.2	6.2	0.933	107	-	15,700	614
16	1-hexadecene	100,000	2.0	4.6	8.5	0.919	88	-104/-10	5581	674
17	1-hexadecene	95,000	2.0	6.0	12.8	0.904	-	-105/ -05		866
18	1-octadecene	61,000	2.1	4.8	3.6	0.940	16, 79	-100/ -58		_
19	1-octadecene	84,000	2.0	5.6	6.2	0.920				
20	1-octadecene	80,000	1.9	7.2	9.4	0.883	99	-92/ -30	135	850
21	1-octadecene	83,000/ 153,000	2.0	11*	14.8	0.86	29	-	-	-
		,,,,,,,								

First figure from GPC-Differential Refractive Index (DRI) with polyethylene standard; second figure from GPC-viscometer.

The gel permeation chromatography (GPC) data for the present copolymer is very unusual in that the Mw as 5 determined by GPC with differential refractive index (DRI) measurement yielded artificially low results as compared to the more accurate (but more difficult) viscosity (VIS) This is apparently due to the length of the measurements. comonomer side chain distributed throughout the polymer backbone. A comparison of calibration curves for converting GPC/DRI data to GPC/VIS developed from the examples is illustrated in Figs. 10 and 11 for dodecene, tetradecene and octadecene copolymers. Standard calibration curves included

10

15

20

25

30

35

in Figs. 10 and 11 for polyethylene, ethylene-propylene copolymer, and ethylene-butene and -hexene copolymers, show the comparatively dramatic differences in the GPC calibration curves for the present copolymers.

The melting point data for some of the examples are with relatively high surprising, particularly those comonomer content. In Examples 10, 13, 14, 18 and 21, note the melting points reflect the crystallinity of the side chains, to the exclusion of the backbone or main chain. Where the side chains introduced by the α -olefin comonomer are frequent enough, usually above about 10 mole percent (or of comonomer less as the length the crystallization of the side chains is evidenced in the lower and/or dual melting points.

The stress-strain properties of the copolymers as reported in Table 3 show that the copolymers are extremely soft and tough materials. The modulus of elasticity can vary from extremely low (note Examples 8 and 20) to moderate, and appears to correlate with both comonomer length and content. The strain to break is very unusual in that it is remarkably high. The strain to break of Examples 17 and 20, in excess of 800 %, is exceptional.

The unusual characteristics of the present copolymers are also seen in the storage modulus (G'), loss modulus (G") and tan δ data developed for Examples 5, 9 and 14 presented below in Table 4. The copolymers in general show that they are very lossy, capable of dissipating substantial energy. This property is very desirable in energy absorption and damping applications, for example, in shock absorbers, vibration dampening, etc. the materials Also, themselves to have good debonding characteristics (G") at high and low frequencies, e.g. they are lossy and compliant, and yet have excellent bonding characteristics (G') at low frequency, for good adhesion performance. See also Fig. 8 which compares $tan \delta$ of the present copolymers against ethylene-butene copolymers at varying comonomer contents.

		goz		268	***	280	82	317	0.41	129	15.6	0.12	74.5	8.87	0.19	48.1	7.52	0.16	1 27	8.07	0.21	32.2	16.6	0.30	675,000	0	0	118	6.74	0.41	45.5	4.05	1.0	45.2	6.86	0.15
		200		229	-	0.30	299	546	0.37	127	15.0	0.12	7.07	8.88	0,13	46.7	8.4	0.18	39.0	9.38	0.24	28.8	10.0	0.35	174,000	3820	0.02	125	7.99	0.33	56.8	8.53	0.15	42.8	69.9	0.17
		901		CZ4	177	0.35	909	196	0.32	122	13.2	0.11	67.2	9.14	0.14	43.5	8.83	0.20	35.3	9.90	0.28	25.1	10.1	15.0	73,200	1290	0.03	108	45.6	0.42	53.7	6.35	0.12	40.7	6.93	0.17
		39.8	Ę	350	138	0.43	559	152	0.27	116	11.9	0.10	62.3	9.53	0.15	38.9	9.56	0.25	29.9	10.2	0.34	19.9	9.88	0.50	46,000	1310	0.03	91.9	27.3	0.30	9.05	5.63	0.11	38.1	7.41	0.19
	FREQUENCY (rad/sec)	0.01	800	077	103	0.45	489	103	0.21	107	11.5	0.11	54.2	10.2	0.19	30.6	10.1	0.33	21.4	9.88	95.0	12.2	8.24	0.68	44,800	1200	0.03	79.0	13.6	0.17	1.72	2.54	0.12	32.5	8.23	0.25
	FREQUENCY	2.51	700		9.68	0.44	434	77.7	0.18	96.9	12.2	0.13	45.4	10.4	0.23	22.0	9.71	0.44	13.4	8.31	0.62	6.23	5.69	0.91	43,400	926	0.05	75.1	7.07	0.09	42.1	6.90	0.16	6*52	8.69	0.34
1		1.00	201		87.8	0.44	407	67.7	0.17	20.7	12.1	0.13	39.6	10.3	0.26	16.5	8.80	0,53	9.00	6.74	0.75	3.62	3.96	1.10	42,900	916	0.05	71.8	6.08	0.08	38.1	7.37	0.19	20.9	8.57	0.41
		0.40	197		8.06	95.0	373	8.09	0.10	65.5	12.4	0.15	33.3	9.80	0.35	11.7	7.74	75.0	5.60	5.05	0.00	1.94	2.55	1.52	45,000	283	0.02	66.9	5.63	0.00	33.6	7.98	0.24	16.1	8.05	0.50
		0,20	197		4.49	0.48	357	28.1	0.10	, 0	12.4	90.10	27.7	9.07	15.0	8.72	6.23	0.72	3.87	3.84	0.99	1.14	1.73	1.52	41,400	(23	0.02	62.1	6.30	0.10	30.7	8.39	0.27	12.7	7.31	0.57
		0.10	196		99.8	0.51	342	2000	0.10	(3.0	<u> </u>	9 0	24.7	8.24	0.35	6.13	5.21	0.85	2.51	2.81	1.12	0.51/	51.15	2.53	40,400	1250	20.0	59.2	6.88	0.12	26.3	8.06	0.31	9.60	6.72	0.70
			6' (dyne x10 ⁵ /cπ ²)	G" (dyne χ1η ⁵ /cm ²)	200	19/119	5		.07.0	5 5		6.76				-5		6"/6"	5	:5	.9/u5	.9	64	6.76°	اد		6.76.	0	ng.	0,0/0,0	ق	ē	,9/ _" 9	.0	5	9/19
	3 E		3,45-			2002	3.00		3,00	,		25.00	, ,		,500	2,64			3,50		28,30	22.5		42000	7 021		3005	-30.0			00ء			25°C		
	STRAIN (%)		-			ç	2		10	2		1	2		٩	2			2		Ş	2		30 0	6.6		٩	2			2			2		
	POLY-	١																						ů												

TABLE 4

-
_
Z
8
C
4
ABLE
=
œ
<

POLY- Mer	STRAIN (%)	TEMP (°C)						FREQUEN	FREQUENCY (rad/sec)				
				0.10	07.0	0,.0	1.00	2.51	10.0	39.8	9	200	202
	10	2°52	9	4.75	6.44	8.97	13.1	17.0	25.3	32.3	36.2	26.1	
			119	4.15	5.27	6.29	7.48	8.32	8.7	8.27	7.74	7 35	7.17
			19/19	0.87	0.82	0.70	0.57	0.47	0.34	0.26	0.21	0, 10	71.0
	10	65°C	19	2.05	2.89	07.7	7.13	10.7	17.1	23.9	28.0	30.0	7.72
			6 "	2.28	3.12	4.10	5.47	6.70	7.90	8.09	7.76	7.43	7,18
			6"/6"	1.11	1.08	0.93	0.77	0.63	97.0	0.34	0.28	70 0	2
Ex. 14	10	ວ•0	6.	0.607	0.959	2.54	4.08	4.08	7.19	11.1	14.0	16.1	18.6
			9	0.793	1.17	2.22	3.00	4.00	4.00	4.72	5.21	5.57	70.5
			G"/G'	1.31	1.22	1.05	0.73	0.56	0.56	0.42	0.37	0.37	25.0
	2	23°C	6,	0.120	0.197	0.319	0.608	1.23	2.58	5.09	7.35	9.23	11.2
			6"	0.198	0.310	0.477	0.813	1.31	2.33	3.50	4.21	4.71	5.11
			12/13	1 45	1 57	4	1.5	7 .	3				

a-tan(delta)

Rheological testing was undertaken for several of the copolymer examples. A relationship of the shear thinning property of the present invention copolymers against comonomer chain length and temperature can be seen in Figs. 3-7. Shear thinning, as represented by the reduction of viscosity (η) and stress (σ) at increasing frequency, is more pronounced for copolymers with shorter comonomer branches $(C_{12}$ versus C_{18} and at a lower temperature). Compare particularly Fig. 4 and Fig. 6.

10

15

20

25

30

35

ADHESIVE FORMULATIONS

Measurements of viscoelastic properties were performed using a PHEOMETRICS SYSTEM IV rheometer or a POLYMER LABORATORIES DMTA rheometer. Isothermal measurements were performed on the SYSTEM IV rheometer over a wide range of temperatures. Isochronal experiments were conducted at a frequency of 10 rad/s and 1 Hz on the SYSTEM IV and the DMTA rheometer, respectively.

Adhesive tests were performed on adhesive compositions dissolved in toluene and then knife-coated to a thickness of about 1.5 mil on a MYLAR substrate. Unless otherwise mentioned, the substrate for the adhesive test was either aluminum, polyethylene or polypropylene.

To prepare a test sample, the adhesive composition was dissolved in toluene and poured inside a hollow-glass cylinder over a piece of stretched cellophane. Films were formed by evaporating the solvent at room temperature. Further drying was conducted in a vacuum oven at 50°C or at room temperature.

The storage modulus (G') is determined according to a Polymer Laboratories, Inc. dynamic mechanical thermal analyzer (DMTA) procedures at ambient temperature. The PSA is cast in a Teflon-coated mold, and 12 mm diameter disks are die cut for DMTA testing. G' is understood in the art to be a measurement of the elastic or storage modulus (stress/strain) measured in phase with sinusoidal shear displacement of the material.

For T-peel testing, the molten adhesive was poured onto a silicone coated release paper and smoothed to a thickness

15

20

of about 6 mils by drawing a heated bar across the adhesive layer. The adhesive film, after cooling was peeled from the release paper and bonded between 2 pieces of 5 mil thick aluminum sheets under the bonding conditions of 150°C/40 psi/10 seconds. T-peel strength is defined as the average load per unit width of bondline required to produce progressive separation of 2 bonded adherends. The separation speed was 2 inches/minute.

The shear adhesion failure temperature (SAFT) was measured as the failure temperature of a tape, coated with a 1.5 mil thickness of the adhesive specimen and adhered on a 1" x 1" overlap onto a steel substrate, under a 500 g vertical load. This test was conducted in an oven by increasing the oven temperature at the rate of 40°F per hour.

EXAMPLES 22-23

Adhesive formulations were prepared by solvent blending in toluene the Example 3 (ethylene- C_{10}) or Example 17 (ethylene- C_{16}) polymer with ESC-5380 tackifier having a hydrogenated cyclic composition ($T_g = 36^{\circ}\text{C}$, $M_{\mu} = 590$ and $M_{\mu}/M_{n} = 1.5$). The formulation was a 60:40 polymer:tackifier weight ratio and either BHT or IRGANOX 1010 was added as a stabilizer (1 wt % based on total adhesive weight).

Adhesive properties are summarized in Table 4. The 25 SAFT is good but the T-Peel results are extraordinary especially for the PE substrate which failed prior to the adhesive bond.

PCT/US92/10590

10

46 TABLE 5

	EXAMPLE 22	EXAMPLE 23
Polymer	Example 3 (C ₂ -C ₁₀)	Example 17 (C ₂ - C ₁₆)
Density (g/ml)	0.94	0.94
MW	123,000	94,000
MP (°C)		118
	T-PEEL (psi)	
Al	5.1	2.25
PE	16.2ª	13.5 ^a
PP	11.3	1.9
	SAFT ^b (°C)	
Al	85	133
PE	94	109

a - substrate failure
b - 1" x 1" x 500g

Differential scanning calorimetry analysis of the Example 23 adhesive (Fig. 1) gives a glass transition temperature (T_g) of 38.4°C. When the ESC-5380 tackifier was replaced in the adhesive blend with ESC-1310LC tackifier (aliphatic composition, $T_g=40$ °C, $M_u=1500$ and $M_u/M_n=1.3$) maintaining the 60:40 polymer:tackifier composition, the T_g was 43.8°C (Fig. 2).

Many modifications and variations besides the embodiments specifically mentioned may be made in the compositions and methods described herein without substantially departing from the concept of the present invention. Accordingly, it should be clearly understood that the form of the invention described herein is exemplary only, and is not intended as a limitation of the scope thereof.

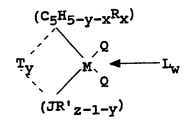
Claims:

- 1. A substantially compositionally uniform copolymer, comprising ethylene and from about 1 to about 50 mole percent of an α-olefin comonomer of at least 10 carbon atoms substantially uniformly incorporated randomly in the copolymer, wherein the copolymer has a density from about 0.85 to about 0.95 g/cm³, a weight average molecular weight from about 30,000 to about 1,000,000 daltons, and a molecular weight distribution from about 2 to about 4.
- 2. The copolymer of claim 1 having a composition distribution breadth index of at least about 70 percent.
- 3. The copolymer of claim 1, wherein the α -olefin comonomer has from 12 to about 100 carbon atoms.
- 4. The copolymer of claim 1, wherein the α -olefin comonomer is straight-chained.
- 5. The copolymer of claim 1, wherein the α -olefin comonomer has from 12 to 30 carbon atoms.
- 6. The copolymer of claim 1, comprising from about 2 to about 30 mole percent of the α -olefin comonomer.
- 7. The copolymer of claim 1, comprising from about 4 to about 30 mole percent of the α -olefin.
- 8. An amorphous copolymer of claim 1, comprising at least about 12 mole percent of the α -olefin comonomer and having a density from about 0.85 to about 0.90 g/cm³.
- 9. A semicrystalline copolymer of claim 1, comprising up to about 12 mole percent of the α -olefin comonomer and having a density above about 0.88 g/cm³.
- 10. The copolymer of claim 1, wherein the molecular weight is from about 80,000 to about 500,000 daltons.
- 11. A film, comprising a semicrystalline copolymer of ethylene and from about 2 to about 12 mole percent of an α-olefin comonomer of at least 10 carbon atoms substantially uniformly incorporated randomly in the copolymer, wherein the copolymer has a density from about 0.88 to about 0.93 g/cm³, a weight average molecular weight from about 80,000 to about 500,000

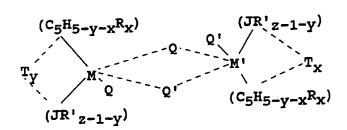
daltons, and a molecular weight distribution from about 2 to about 4.

- 12. An adhesive, comprising the copolymer of claim 8 blended with a tackifier.
- 13. A method for preparing a substantially compositionally uniform copolymer of ethylene and an α -olefin comonomer having at least 10 carbon atoms, comprising contacting a mixture of ethylene and the comonomer with a catalyst at polymerization conditions wherein the ethylene:comonomer reactivity ratio is less than about 50.
- 14. The method of claim 13, wherein the reactivity ratio is less than about 30.
- 15. The method of claim 13, wherein the reactivity ratio is from about 3 to about 20.
- 16. The method of claim 13, wherein the reactivity ratio is from about 5 to about 15.
- 17. A method for preparing the copolymer of claim 1, comprising:

charging a reactor with ethylene and the α -olefin at reaction conditions in the presence of a catalyst system including an activated metallocene catalyst component of the formula:



or



wherein M is Zr, Hf or Ti in its highest formal oxidation state:

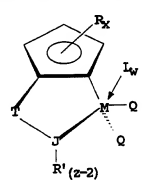
 $(C_5H_{5-\gamma-x}R_x)$ is a cyclopentadienyl ring which is with from zero to substituted substituent groups R, "x" is 0, 1, 2, 3, 4 or 5 denoting the degree of substitution, and each substituent group R is, independently, a radical selected from a group consisting of C_1-C_{20} hydrocarbyl radicals; substituted C_1-C_{20} hydrocarbyl radicals wherein one or hydrogen atoms is replaced by a halogen phosphido amido radical, a radical, an radical, an alkoxy radical, an alkylborido radical or a radical containing a Lewis functionality; C1-C20 basic acidic or radicals hydrocarbyl-substituted metalloid wherein the metalloid is selected from the Group IV A of the Periodic Table of Elements; amido radicals, halogen radicals, and radicals, alkoxy radicals. phosphido alkylborido radicals, or a radical containing Lewis acidic or basic functionality; or (C5H5. v-rRr) is a cyclopentadienyl ring in which two adjacent R-groups are joined forming C4-C20 ring to give a saturated or unsaturated polycyclic cyclopentadienyl ligand;

(JR',-|-v) is a heteroatom ligand in which J is an element with a coordination number of three element with from Group V A or an coordination number of two from Group VI A of the Periodic Table of Elements, each R' is, independently a radical selected from a group consisting of C1-C20 hydrocarbyl radicals, C1-C20 hydrocarbyl radicals substituted or more hydrogen wherein one atoms replaced by a halogen radical, an amido radical, an alkylborido radical, a phosphido radical, an alkoxy radical, or a radical acidic containing а Lewis or basic

- functionality; and "z" is the coordination
 number of the element J;
- each Q is, independently, any univalent anionic ligand, provided that where Q is a hydrocarbyl such Q is different than the (C₅H_{5-y-x}R_x) or both Q together are an alkylidene, a cyclometallated hydrocarbyl or a divalent anionic chelating ligand;
- "y" is 0 or 1 when "w" is greater than 0; "y" is 1 when "w" is 0; when "y" is 1, T is a covalent bridging group containing a Group IV A or V A element;
- L is a neutral Lewis base where "w" denotes a number from 0 to 3.
- 18. The method of claim 17, wherein the catalyst system includes an alumoxane component as the metallocene activator.
- 19. The method of claim 17, wherein the metallocene catalyst component contains at least one substituent-capable of reacting with a proton and the catalyst system comprises the metallocene catalyst component activated by combination with a cation capable of donating a proton and a bulky non-coordinating anion capable of stabilizing the metal cation formed as a result of reaction between the proton provided by the cation and said substituent of the metallocene.
- 20. The method of claim 17, wherein M is Ti in its highest formal oxidation state.
- 21. The method of claim 17, wherein the heteroatom ligand group J element is nitrogen, phosphorous, oxygen or sulfur.
- 22. The method of claim 21, wherein Q is a halogen or hydrocarbyl radical.
- 23. The method of claim 21, wherein the heteroatom ligand group J element is nitrogen.
- 24. The method of claim 21, wherein R is a C_1-C_{20} hydrocarbyl radical and "x" is 4.
- 25. The method of claim 21, wherein R' is an aliphatic hydrocarbyl radical having either a primary or

secondary carbon atom bonded directly to the J element or an alicyclic hydrocarbyl radical having a secondary carbon atom bonded directly to the J element.

- 26. The method of claim 21, wherein the mole ratio of Al:M is from 0.5:1 to 5000:1.
- 27. The method of claim 21, wherein Q is independently halogen, hydride, or a substituted or unsubstituted C_1 - C_{20} hydrocarbyl, alkoxide, aryloxide, amide, arylamide, phosphide or aryl phosphide, provided that where any Q is a hydrocarbyl such Q is different from $(C_5H_{4-x}R_x)$ or both together are an alkylidene or a cyclometallated hydrocarbyl.
- 28. The method of claim 18, wherein said alumoxane compound is methylalumoxane.
- 29. The method of claim 21, wherein the catalyst system includes an alumoxane and the Group IV B transition metal component has the formula:

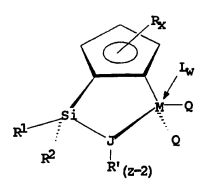


wherein M is Zr, Hf or Ti in its highest formal oxidation state;

R is a substituent group with "x" denoting the degree of substitution (x = 0, 1, 2, 3 or 4) and each R is, independently, a radical selected from a group consisting of C_1 - C_{20} hydrocarbyl radicals, substituted C_1 - C_{20} hydrocarbyl radicals wherein one or more hydrogen atoms is replaced by a halogen radical, an amido radical, a phosphido radical, an alkoxy radical or any other radical containing a Lewis acidic or basic functionality, C_1 - C_{20} hydrocarbyl-substituted

metalloid radicals wherein the metalloid is selected from the Group IV A of the Periodic Table of Elements and halogen radicals, amido radicals, radicals, phosphido radicals, alkylborido radicals or a radical basic : Lewis acidic containing functionality, or at least two adjacent Rgroups are joined forming C_4 - C_{20} ring to give unsaturated polycyclic saturated or cyclopentadienyl ligand;

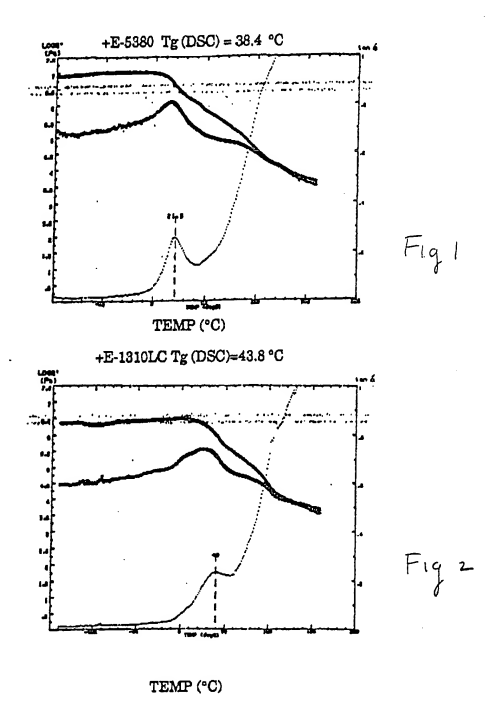
- (JR'_{z-2}) is a heteroatom ligand in which J is an element with a coordination number of three from Group V A or an element with coordination number of two from Group IV A or the Periodic Table of Elements, and each R' is, independently a radical selected from a C1-C20 hydrocarbyl consisting of group hydrocarbyl C1-C20 radicals, substituted radicals where one or more hydrogen atom is replaced by a halogen radical, an amido radical, a phosphido radical, and alkoxy radical or a radical containing a Lewis acidic or basic functionality, and "z" is the coordination number of the element J;
- each Q is, independently, any univalent anionic ligand, such as a halide, hydride, or a substituted or unsubstituted C_1 - C_{20} hydrocarbyl, alkoxide, aryloxide, amide, arylamide, phosphide or arylphosphide, or both Q together are an alkylidene, or a cyclometallated hydrocarbyl or any divalent anionic chelating ligand;
- T is a covalent bridging group containing a Group VI A or V A element;
- L is a neutral Lewis base where "w" denotes a number from 0 to 3.
- 30. The method of claim 21, wherein the catalyst system includes an alumoxane and the Group IV B transition metal component has the formula



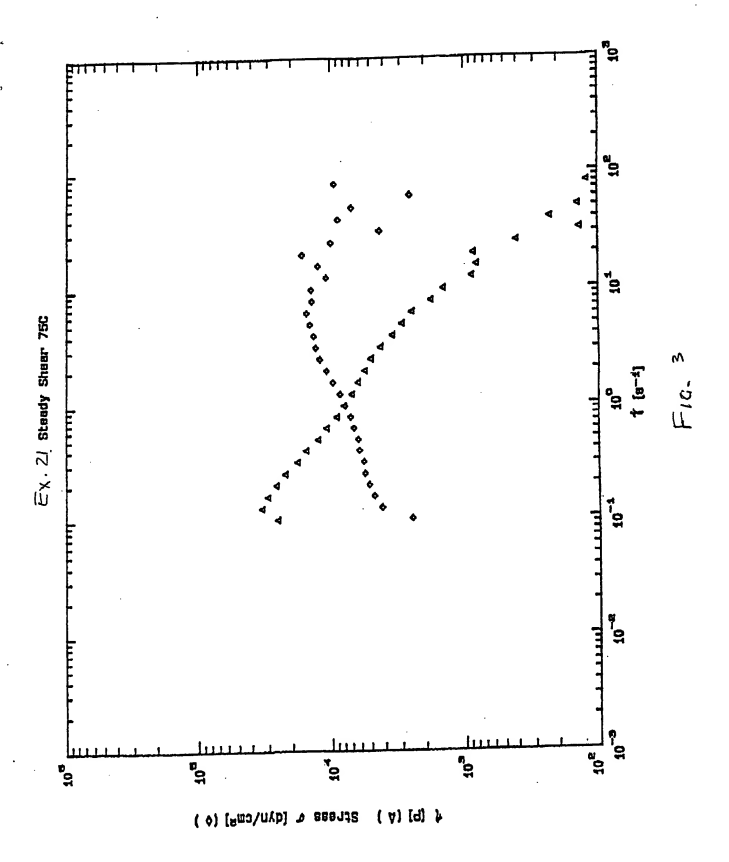
wherein R^1 and R^2 are, independently, C_1 to C_{20} hydrocarbyl radicals, substituted C_1 to C_{20} hydrocarbyl radicals wherein one or more hydrogen atoms is replaced by a halogen atom; R_1 and R_2 may also be joined forming a C_3 to C_{20} ring.

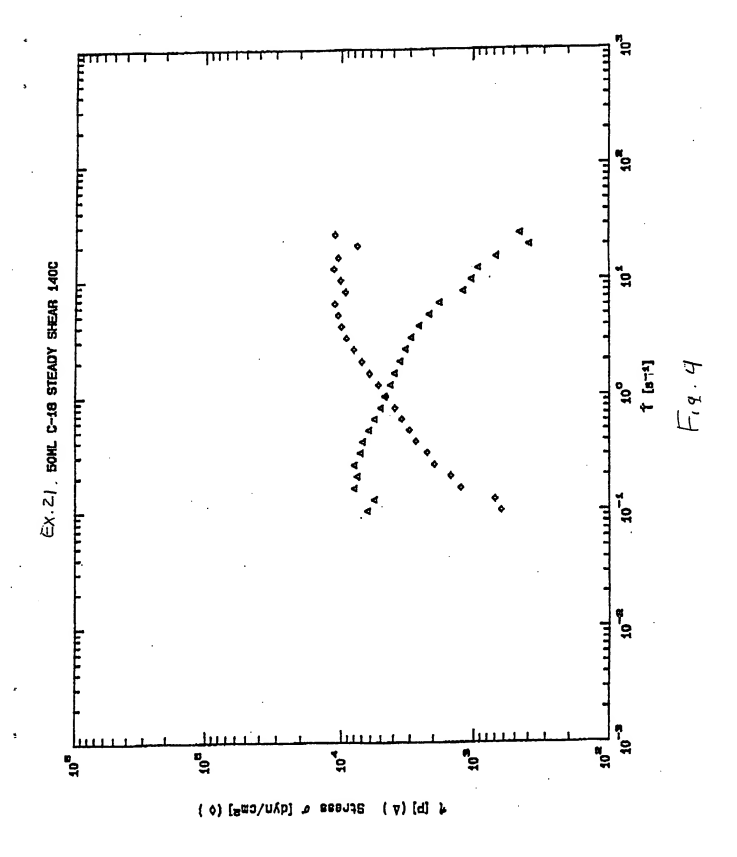
- 31. The method of claims 29 or 30, wherein J is nitrogen.
- 32. The method of claim 31 wherein R is a C₁ to C₂₀ hydrocarbyl radical, "x" is 4 and R' is an aliphatic hydrocarbyl radical having either a primary or secondary carbon atom bonded directly to the J element or an alicyclic hydrocarbyl radical having a secondary carbon atom bonded directly to the J element.
- 33. The method of claims 29 or 30, wherein M is titanium.
- 34. The method of claims 29 or 30, wherein M is hafnium or zirconium.
- 35. The method of claim 29, wherein T is silicon, J is nitrogen and when R is an alkyl radical, R' is a cyclohydrocarbyl, and when "x" is 4 and the R substituents form a polycyclic ring system, R' is an alkyl or cyclohydrocarbyl radical.
- 36. The method of claim 30, wherein M is titanium, J is nitrogen and R' is cyclohydrocarbyl.
- 37. the method of claim 36, wherein R' has from 10 to 16 carbon atoms.
- 38. The method of claim 36, wherein R' is cyclododecyl.
- 39. The method of claim 18, 29 or 30, wherein the Al to transition metal molar ratio is 2000:1 or less.
- 40. The method of claim 17, wherein the comonomer has from 12 to 30 carbon atoms.
- 41. The method of claim 17, wherein the comonomer is straight-chained.

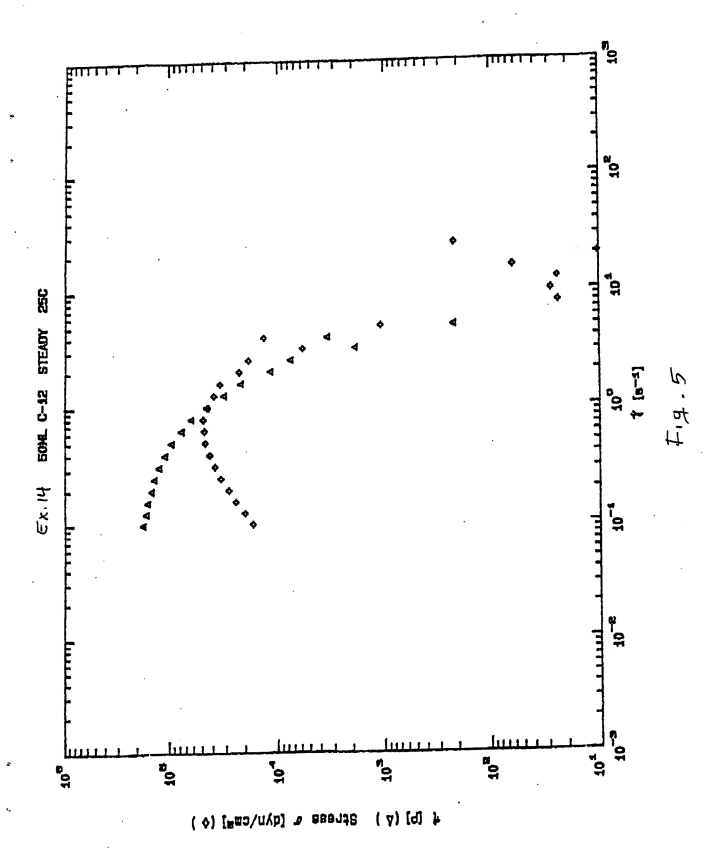
(C2/C16) COPOLYMER, TACKIFIER BLENDS (60/40 WT/WT)

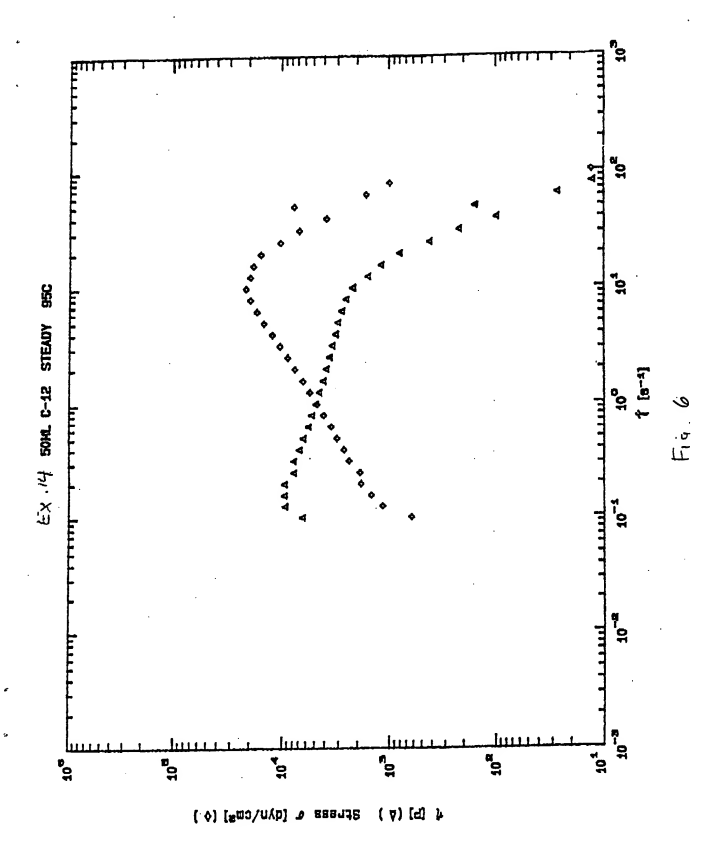


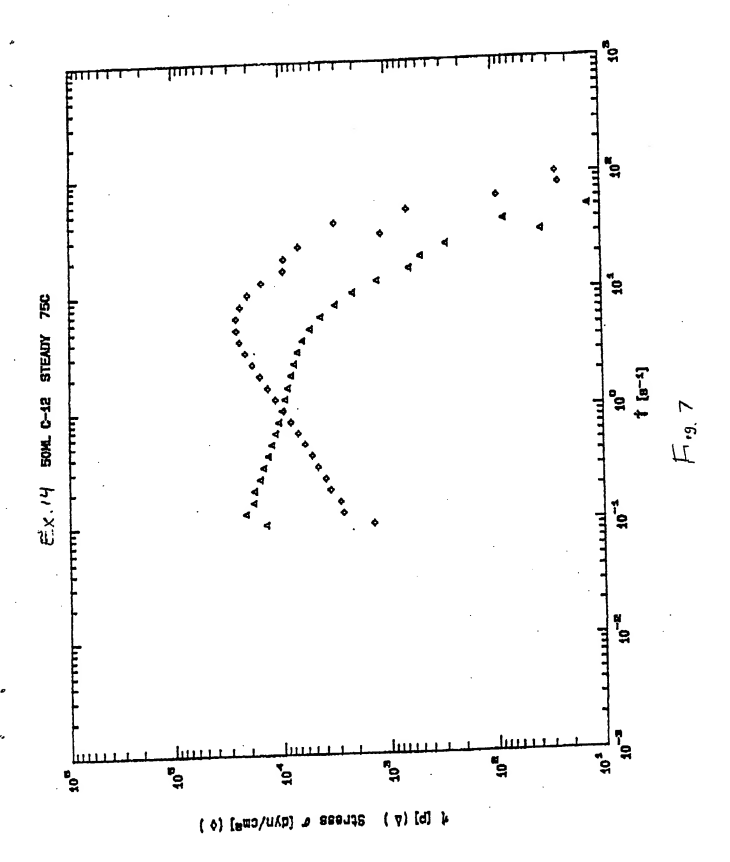
C21/C16 Contains 6.0 mole % comonomer MN = 94K MND = 2.0 Density = 0.904 g/cc



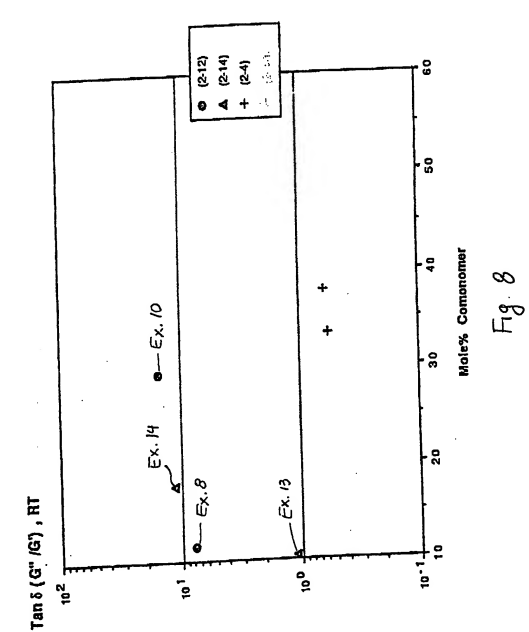






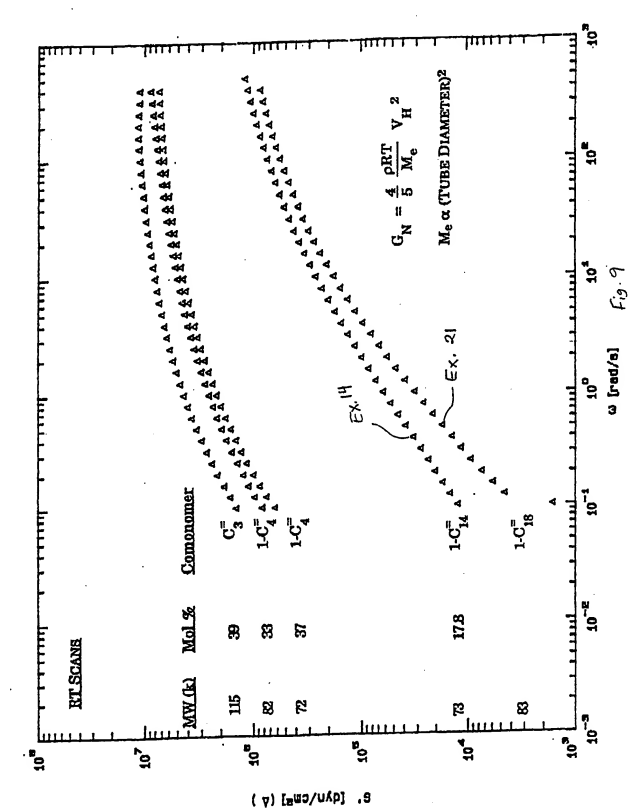


Tan 8 (G"/G')a of Ethylene Copolymers Increases with Branch Length



a G' @ 100 Rad/Sec, G' @ 1 Rad/Sec; 10% Strain; Rheometrics System IV.

Effect of Branch Length on Storage Modulus Ethylene Copolymers



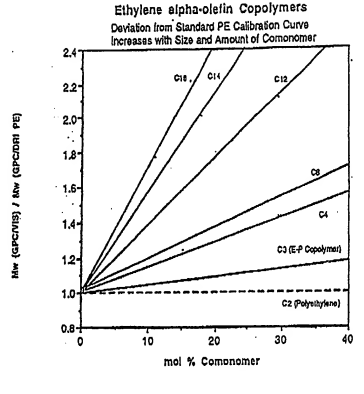
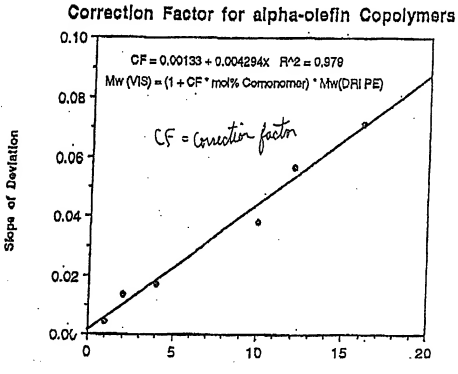


Fig. 10



Number of Carbon Atoms in Side Chains

Fig. 11

INTERNATIONAL SEARCH REPORT

International Applica No PCT/US 92/10590

	TO TON OR SIDE	CT MATTER (If several classification sym	bols apply, indicate all	1)6	
According Int.Cl	to International Patent . 5	Classification (IPC) or to both National Class C 08 F 210/02 C 08	sification and IPC	C 08 F 2Ï0/	116
II. FIELDS	SEARCHED		ii - Samball		
		Minimum Document			
Classificat	ion System	Cla	assification Symbols		
Int.Cl	.5	C 08 F			
		Documentation Searched other the to the Extent that such Documents are	an Minimum Docume Included in the Field	ntation s Searched ⁸	
III. DOCUI	MENTS CONSIDERE	D TO BE RELEVANT 9			Relevant to Claim No.13
Category °	Citation of De	ocument, 11 with indication, where appropriate	e, of the relevant pass	ages 17	Refevant to Clarin 110.
Х		007526 (MITSUI) 12 July aim 1; page 17, lines 12	, 1990		1
х	FP A N	420436 (EXXON CHEM.) 3 see claims; page 9, line	April		16,1
x	EP,A,0 1991,	436520 (GAS RESEARCH) I see the whole document	lO July		1
X	EP,A,0 13 Nov	161060 (NORTHERN PETROC ember 1985, see claims	CHEM.) -/-		1
"A" do "E" ea fli "L" do wh cit	nsidered to be of partic rhier document but pub ing date coment which may thro ich is cited to establish ation or other special t	meral state of the art which is not miar relevance lished on or after the international ow doubts on priority claim(s) or the publication date of another	or priority date cited to underst invention "X" document of pa cannot be consilinvoive an inve document of pa cannot be considocument is coments, such coments, such consiling to the constant of the cannot be consiling to the constant is coments, such consiling to the constant of the cannot be consiling to the cannot be consiling to the constant of the cannot be consiling to the cannot be cannot be consiling to the cannot be cannot be consiling to the cannot be	published after the internand not in conflict with the principle or theoretical relevance; the cial dered novel or cannot be nive step ritcular relevance; the cial dered to involve an inventional with one or more mbination being obvious	ry underlying the imed invention considered to timed invention tive step when the other such docu-
P do	cument published prior ter than the priority da	to the international filing date but te claimed	in the art. "&" document mem	ber of the same patent fa	mily
	IFICATION		Deep of Mailine	of this International Sec	urch Report
Date of the	Actual Completion of 08-02-	the International Search 1993		1 9. 03. 93	
<u> </u>			Signature of Au	nthorized Officer	
Internation	al Searching Authority EUROPI	CAN PATENT OFFICE		DE ROECK	

Page 2
International Applicat io PCT/US 92/10590

(CONTINUED FROM THE SECOND SHEET) III. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to Claim No. Citation of Document, with Indication, where appropriate, of the relevant passages Category o WO,A,8500172 (NESTE OY) 17 January 1985, see claims; examples 1 A

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9210590

SA 68059

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 23/02/93

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent document cited in search report	Publication date		nt family nber(s)	Publication date
WO-A- 9007526	12-07-90	JP-A- JP-A- JP-A- EP-A- JP-A-	2173014 2173016 2173015 0495099 2276807	04-07-90 04-07-90 04-07-90 22-07-92 13-11-90
EP-A- 0420436	03-04-91	US-A- AU-A- CA-A- CA-A- EP-A- JP-A- WO-A- US-A- US-A- US-A-	5055438 6248390 6443990 2024899 2065745 0491842 3188092 9104257 5026798 5057475 5168111 5096867	08-10-91 21-03-91 18-04-91 14-03-91 01-07-92 16-08-91 04-04-91 25-06-91 15-10-91 01-12-92 17-03-92
EP-A- 0436520	10-07-91	None		
EP-A- 0161060	13-11-85	CA-A- JP-A- JP-B- US-A-	1256246 1308410 4023645 4754007	20-06-89 13-12-89 22-04-92 28-06-88
WO-A- 8500172	17-01-85	BE-A- DE-C- DE-T- FR-A- GB-A- JP-T- NL-T- US-A-	899984 3490300 3490300 2549073 2152517 60501609 8420184 4690991	15-10-84 07-05-92 15-05-85 18-01-85 07-08-85 26-09-85 01-04-85